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Improving magnesium's corrosion resistance through tannic acid-polyethyleneimine coatings for bioresorbable implant applications

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The high corrosion rate of magnesium limits its application as a bioresorbable implant, despite its good biocompatibility. A tannic acid (TA) coating on magnesium has succeeded in increasing the corrosion resistance of magnesium, although the presence of cracks in this coating still limits its effectiveness. This research focuses on the effect of adding polyethyleneimine (PEI) to the TA coating to increase the corrosion resistance of magnesium and its biocompatibility. The TA-PEI coating on magnesium with TA: PEI weight ratios of 1:0, 1:0.5, 1:1, and 1:2, which was achieved by the immersion method, was evaluated by characterizing the composition, morphology, structure, and surface roughness of the coating as well as the deformation mechanism. The corrosion properties and corrosion rate were examined through immersion and electrochemical corrosion tests (potentiodynamic polarization and EIS). Wettability, cell adhesion, and MTT assay tests were carried out as a coating biocompatibility test. The magnesium's surface was successfully covered with the TA-PEI coating, increasing its corrosion resistance and biocompatibility. The increase in the PEI content in the weight ratio of TA: PEI from 1:0 to 1:2 resulted in a reduction in the areal crack density from 25.49% to 7.12%, an increase in the coating thickness from 5.7 μm to 9.3 μm, a decrease in the surface roughness from 5.20 μm to 0.98 μm, but a change in the deformation mechanism from ductile to brittle. Adding PEI to TA as a pure magnesium coating has increased the material's corrosion resistance. The changed weight ratio of TA: PEI from 1:0 to 1:2 is the most optimum ratio because it can reduce the corrosion rate of the material from 3.7 mm year<sup>-1</sup> to 0.47 mm year<sup>-1</sup>. The formation of hydroxyapatite on a corroded material indicates that this system is biocompatible. Increasing the PEI content from 1:0 to 1:2 TA: PEI weight ratio decreased the contact angle from 74° to 24°. At a 1:2 TA: PEI ratio, passage 6 HWJ cells can adhere well, and this system has the highest absorbance from the MTT assay result. Both indicate that this coating system has good biocompatibility with human physiochemical systems.

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### 1. Introduction

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Bone is one of the musculoskeletal systems with specific mechanical properties.<sup>1-3</sup> The load-bearing capacity of bone is limited; thus, it will fracture when the load exceeds a certain

<sup>d</sup> Biotechnology Research Group, School of Life Sciences and Technology, Institut Teknologi Bandung, Jl. Ganesha 10, Bandung 40132, Indonesia value. However, the bone itself has the ability to regenerate its structure and functionality within 4 months to 1 year.<sup>2</sup>

To realign fractured bones, orthopaedic implants are used to support the regeneration process. Implants must be biocompatible since they are in contact with the living system inside the human body.<sup>4–6</sup> Metals, *e.g.*, titanium alloy, chromium–cobalt alloy, and stainless steel, have been developed as bioinert materials. However, there are several downsides to using these bioinert materials. Under several operating conditions, the metals will be dissolved and form toxic ions in the body. The stress shielding effect has also been observed due to a significant difference in the stiffness value between bone and implants.<sup>7</sup> Implant patients also need to be operated on twice, insertion and extraction. Thus, a biodegradable material with good biocompatibility and similar stiffness to the bone can be a solution.<sup>8</sup>

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Magnesium (Mg) is a promising biodegradable material since it is abundant in the human body and has a similar stiffness value compared to bone.8,9 Nonetheless, the high degradation rate has limited its use in orthopaedic implants. Premature mechanical loss would likely occur in Mg-based implants before the fractured bone is fully restored.<sup>10</sup> To overcome this problem, alloying and coating of Mg can be employed to hinder corrosion.<sup>11-47</sup> By adding some alloying elements to magnesium at a specific value, the corrosion resistance of magnesium will enhance. In addition, the mechanical properties will also change. However, this process is complex because alloying should be done in a liquid state while magnesium has high partial pressure, making it easier to form a gas state. If the atmosphere is not being controlled carefully with inert gases, the targeted final composition is hard to achieve.<sup>15,21</sup> Compared to alloying Mg, the coating process of pure Mg would be relatively cheap and easily done since pure Mg has sufficient stiffness and strength needed as an orthopaedic implant.48

Tannic acid (TA) is one of the largest natural resources easily found, such as chestnuts and mangroves.<sup>49</sup> TA is a polyphenol molecule consisting of abundant catechol groups with orthohydroxyl groups. These ortho-hydroxyls make TA suitable for reaction with metals through metal-chelation to form TA-metal complexes.<sup>28,50,51</sup> Furthermore, TA is well-known for its biocompatibility because this molecule is hydrolysable, biocompatible, and has superior hemostatic properties due to its ability to react with various proteins.<sup>49,52</sup> For these reasons, including its wide range of availability and the facile deposition method through immersion, TA is well developed as a coating material for Mg to enhance corrosion resistance, known as a conversion coating.<sup>31,53</sup> The effectiveness of the TA coating in overcoming corrosion resistance has been proven compared to others; however, the morphology of the TA coating is full of cracks which enables fluids such as simulated body fluid (SBF) to penetrate the Mg substrate and cause corrosion.28,31

Nevertheless, TA has excellent adhesiveness, which means that TA can provide intermolecular reactions with nucleophiles, such as a molecule that contains amine bonds.<sup>28,31,45,49</sup> Polyethyleneimine (PEI) is a molecule that contains amine groups which means that this molecule can make a linkage with TA. The catechol group will be oxidized and form primary boding with amine groups in PEI, also known as crosslink linkage through Michael-type addition or a Schiff-base reaction.<sup>28,50,54,55</sup> This phenomenon could affect the morphology of the coating, that is, reducing the crack density, which means a more solid and uniform coating to give better corrosion protection, as well as improving other properties such as enhancing wettability and facilitating cell adsorption.<sup>28</sup>

In this research, various TA–PEI coatings were deposited on the Mg substrate, aiming to study the effect of PEI addition on the TA coating and its morphology based on the visual investigation through SEM and corrosion behaviour through immersion and electrochemical tests. Moreover, the biocompatibility based on *in vitro* tests, cell adhesion, and MTT assay was also investigated in this research.

### 2. Materials and methods

#### 2.1. Materials

Pure Mg (with an Mg content of 99.3 wt%) was used as the substrate. TA powder (Merck, Germany) and the PEI oligomer (Jinan Future Chemical Co. Ltd, China;  $M_w = 1300$ ) were used as the coating material. Aqua DM was also used throughout this study as a solvent material.

#### 2.2. Synthesis of TA-PEI coatings on pure Mg

A pure Mg surface was first prepared by grinding and polishing. The coating material was prepared by mixing TA and PEI with weight ratios of TA: PEI of 1:0, 1:0.5, 1:1, and 1:2. Pure Mg was then immersed in the mixture for 5 minutes. After 5 minutes, the coated Mg was dried in ambient air for approximately 1 min. The immersion process was repeated three times in a fresh mixture of TA-PEI at the same time interval. Finally, the coated Mg was dried in ambient air for 24 hours. The sample codification is shown in Table 1.

#### 2.3. Chemical structure and surface characterization

After treatment, Fourier transform infrared (FTIR) spectroscopy was used to analyze the functional groups of the coating (Bruker Alpha II, Bandung, Indonesia). The components and morphology of the surface were examined using a scanning electron microscope (SEM) integrated with an energy dispersive spectroscopy system (EDS) (JSM-IT300, Bandung, Indonesia). The areal crack density of the samples was calculated using ImageJ software by identifying percentage areas with a discontinuous coating morphology. EDS line analysis was used to quantify the coating thickness on the coating's cross-section. The coating's thickness was considered a part of the coated material with abundant elements from the coating without any presence of magnesium elements. The border between the coating and the substrate is defined when the amount of magnesium detected from the EDS line analysis suddenly increases. In addition, the deformation mode was also visually examined by SEM after the micro-Vickers hardness test (0.01 HV) was carried out. The surface of coating roughness was measured through the contact (tactile) method (Mitutoyo, Bandung, Indonesia).

#### 2.4. Immersion corrosion test and corrosion product analysis

One side of all samples was sealed with tape to expose only one side of the sample  $(1 \text{ cm} \times 1 \text{ cm})$  to SBF solution as a corrosive fluid representing the human physiological system. The immersion tests of samples were conducted in a chamber that contained simulated body fluid (SBF) solution with pH 7.4

Table 1	Sample codifica	ation			
Code	TA (mg/500 mL)	PEI (mg/500 mL)	Weigh ratio	Solvent	Dimensions (cm)
TA TA–1PEI TA–2PEI TA–4PEI	2	0 1 2 4	1:0 1:0.5 1:1 1:2	Aqua DM	$1 \times 1 \times 0.2$

and a temperature of  $37 \pm 1$  °C. Samples were immersed with different intervals of immersion (1 hour, 1 day, 3 days, and 7 days) in a 40 mL SBF for each sample to stabilize the pH solution based on ASTM G31-72. Furthermore, SBF is constantly changed, and pH solutions were also measured every 24 hours. The surface morphology after the immersion test was examined using a scanning electron microscope (SEM) (JSM-IT300, Bandung, Indonesia). Corrosion products were analyzed using an XRD machine (Bruker D8 Advance). The percentage of each phase of the corrosion products was analyzed using MATCH! Software.

#### 2.5. Electrochemical corrosion test

One side of all samples was connected to copper wire and sealed with tape to expose only one side of the sample  $(1 \text{ cm} \times 1 \text{ cm})$  to SBF for both potentiodynamic polarization and electro impedance spectroscopy (EIS) test. The potentiodynamic polarization test was conducted with a three electrode configuration. The samples were connected to the working electrode, graphite to the counter electrode, and saturated calomel to the reference electrode. A potentiodynamic polarization test (VersaSTAT3, Bandung, Indonesia) was only conducted on fresh samples. Meanwhile, an EIS test (Gamry, Bandung, Indonesia) with the same configuration of electrodes was performed on samples immersed at SBF for 1 hour, 1 day, 3 days, and 7 days using 5 mV sinusoidal potential with 100 kHz to 0.01 Hz frequency range.

#### 2.6. Water contact angle test

The wettability of the samples was measured using a contact angle test (Dino-Lite, Bandung, Indonesia). Specimens are placed on a flat horizontal surface and subjected to one drop of water through a pipette. The angle formed between the water and specimens was then captured using a camera integrated with DinoCapture 2.0 software. To ensure repeatability, all tests were repeated three times.

#### 2.7. Cell adhesion and MTT assay test

Passage 6 human Wharton's jelly mesenchymal stem cells (HWJ-MSCs) were used in this experiment. DMEM (Dulbecco's Modified Eagle's Medium) with 10% FBS and 1% antibiotic-antimycotic was used to culture the cells. The medium was held at 37  $^{\circ}$ C, with 95% humidity and 5% CO<sub>2</sub>. Furthermore, cell fixation was carried out with 100 µl of 2.5 glutaraldehyde in 0.1 cacodylate buffer, incubated overnight at 4  $^{\circ}$ C. After that, alcohol series and HDMS were used to dehydrate and dry the samples, respectively. After fixation, the cell morphologies were examined by scanning electron microscopy (JSM-IT300, Bandung, Indonesia).

For the MTT assay test, 24-well plates were used as the media. Passage 6 HWJ-MSCs were grown at a density of 15 000 cells per well. The plates were incubated for 1, 3, and 7 days after adding the medium to the well. The temperature should be maintained at 37 °C. Cell viability was assessed using an MTT test with 0.5 mg mL<sup>-1</sup> as the final concentration.

### Results and discussion

# 3.1. Functional group characterization of TA and TA-PEI coated pure Mg

The FTIR spectra of TA-PEI coatings are shown in Fig. 1a. There are several distinct bands observed.

A band at 1710.86 cm<sup>-1</sup> of the TA sample shows the presence of C=O functional groups in the TA structure. Bands at 2960.73–2962.66 cm<sup>-1</sup> were only observed on TA–PEI representing CH<sub>2</sub> functional groups at the PEI structure. This evidence matches the previous study to show the presence of both functional groups.44,45 Broad bands were observed around 3373.50-3421.08 cm<sup>-1</sup>, showing chemical bonding evidence of TA and PEI.28 More detailed information on the chemical bonding of TA-PEI is illustrated in the schematic figure, shown in Fig. 1b. Catechol groups in TA can spontaneously be oxidized into quinone groups when TA remains in touch with an alkaline solution. When PEI is added to the same solution, the oxidized catechol group will react with the amino group in PEI through a Michael-type or Schiff-base reaction to build a primary chemical bonding. In other words, crosslink bonding is formed between TA and PEI. The linkage makes overlap stretching vibration between TA and PEI around 2500-3700 cm<sup>-1</sup>. Thus, individual vibration of the OH structure at TA and NH structures at PEI reduces.<sup>44</sup> The origin bands of OH and NH<sub>2</sub> are at 3392 cm<sup>-1</sup> and 3371 cm<sup>-1</sup>, respectively.<sup>45</sup> However, this study observed TA, TA-1PEI, TA-2PEI, and TA-4PEI bands



Fig. 1 (a) FTIR spectrum of TA-PEI coated Mg. (b) Schematic of the TA and PEI chemical reaction on the pure Mg substrate.

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at 3373.50 cm<sup>-1</sup>, 3412.08 cm<sup>-1</sup>, 3410.15 cm<sup>-1</sup>, and 3392.72 cm<sup>-1</sup>, respectively. This evidence shows a band shift due to the strong hydrogen bonding between TA and PEI besides the presence of crosslink phenomena.<sup>45</sup> In addition, there are also metal chelation phenomena between TA and metal at the material surface. This type of bonding is not strong enough as TA–PEI bonding but enough to make the coating deposited and bonded on a metal surface.

# 3.2. Surface characterization of TA and TA-PEI coated pure Mg

**3.2.1. Surface and cross-sectional morphology.** Fig. 2a shows the top-view side morphology of TA and TA-PEI-coated pure Mg. It was shown that the coatings had a discontinuous morphology. From a two-dimensional approach, the areal crack density was defined as a percentage of discontinuous coating parts from the top view of the coating. Based on visual examination of the SEM figure, Fig. 2a and b, and crack density calculation, it was found that the higher the addition of PEI, the lower the number of cracks due to crosslinking between TA and PEI. Thus, the coating tends to be more uniform and continuous. Furthermore, the surface roughness resulting from macro testing on the whole coating's surface area also supports the coating structure on the micrometer scale. With the increase of PEI, the surface roughness of the coating is reduced

due to the lower crack density formed. Quantitatively, the increase in the PEI content from a weight ratio of TA: PEI 1:0 to 1:2 resulted in a reduction in the area of crack density from 25.49% to 7.12%, and a decrease in the surface roughness from 5.20  $\mu$ m to 0.98  $\mu$ m.

In addition, the EDS result on all coatings shows evidence of Mg, C, and O as an element that builds TA, PEI, and Mg as a substrate. Meanwhile, N was only observed on TA-PEI samples due to the presence of  $NH_2$  in PEI, see Table 2.

From a cross-sectional view of the coatings shown in Fig. 2c, the morphology of coatings is uneven and not continuous due to the presence of cracks, confirming the morphology from the top view. It can be seen that the crack penetrated through the thickness of the coating until it reached the substrate. Based on the EDS line analysis result in Fig. 2d, the coating tends to be thicker with the increase of PEI addition due to the higher cross-link density between the TA and PEI. Thus, the deposition of TA-PEI is much more to occur and be deposited on magnesium. EDS line analysis result shows that the increase in the PEI content from a weight ratio of TA : PEI 1:0 to 1:2 resulted in an increase in the coating thickness from 5.7  $\mu$ m to 9.3  $\mu$ m. A simple mathematical equation with linear behaviour assumptions shows that adding 1 mg/500 mL of PEI will increase the coating thickness to 0.95  $\mu$ m.

**3.2.2. Deformation behaviour of coating.** The deformation behaviour of the coating under a specific mechanical load is



Fig. 2 (a) Surface morphology of TA and TA–PEI coated pure Mg, observed through SEM (500×). (b) Coating's crack density. (c) Cross-sectional view of TA and TA–PEI coated pure Mg. (d) Coating thickness, observed through SEM (3000×).

 Table 2
 EDS result of all coated samples

Samples	Elements (wt%)					
	С	Ν	0	Mg		
ТА	31.21	0	36.55	32.24		
TA-1PEI	38.35	3.08	30.24	28.15		
TA-2PEI	47.94	3.95	15.29	32.82		
TA-4PEI	35.07	3.82	28.32	32.79		

shown in Fig. 3. In this section, the fracture of the polymeric coating will use fibrous and cleavage terms. Although this term is widely used in metallic materials, some researchers still use this to define fracture behaviour in polymeric materials.<sup>56-59</sup> Furthermore, although deformation behaviours in this coating were examined on the coating's surface (not in bulky specimens), this characterization is still suitable to predict the fracture behaviour of the polymer because fracture related to the strength of the polymers still correlates with the localized deformation behaviour of the material's surface.<sup>60</sup> TA samples show fibrous characteristics, while TA-PEI shows cleavage characteristics. Without the crosslinking phenomenon, the polymer chain is flexible and can efficiently distribute the mechanical load, and that is why TA can plastically deform before fracture.<sup>58</sup> Meanwhile, with the addition of a crosslinker agent, in this case, PEI, the polymer chain tends to be more rigid. If the chain is rigid, it would be difficult to distribute the mechanical load and plastically deform before fracture. The polymer chain will absorb the energy without distributing it to the other chains, and finally, a brittle fracture occurs.

## 3.3. Immersion corrosion behaviour of TA and TA-PEI coated pure Mg

**3.3.1. pH variation.** As shown in Fig. 4, pH variation occurred during the immersion test. On the first day of immersion, the pH values of all samples increase rapidly, then become constant and stabilize at 8.1 after 3 days of immersion. The pH values did not change until the 7th day of immersion. SBF can penetrate the substrate through the through-the-thickness cracks of the coating as a corrosive fluid. After corrosive fluid contact with the substrate,



the electrochemical reaction occurred due to the formation of the cathodic and anodic sites of the substrate. The cathodic reaction will generate OH<sup>-</sup>, which is why the pH increases.<sup>15,19</sup> Upon increasing the immersion time, corrosion products will form and deposit on coating surfaces, which impede SBF penetration into the substrate. Consequently, the area exposed to SBF will be reduced, and the electrochemical reaction will be diminished. Thus, the pH values become stable.

**3.3.2.** Surface morphology after the immersion test and its corrosion products. The surface morphology of all samples after the immersion test for 7 days is shown in Fig. 5. Corrosion products were deposited on all surfaces of all samples. Corrosion products are deposited not only on the continuous part of the coating but also in the crack area. Thus, this will reduce the observed areal crack density for each sample after the immersion time, see Fig. 5b.

Table 3 shows the EDS result of the corrosion product. The primary corrosion product elements are magnesium, oxygen, calcium, and phosphor. It can be predicted that brucite or magnesium hydroxide  $(Mg(OH)_2)$  and hydroxyapatite



Fig. 3 Deformation behaviour of TA and TA-PEI coated pure Mg under mechanical load.



Fig. 5 Surface morphology of TA and TA-PEI coated pure Mg after immersion for 7 days.

Table 3 EDS analysis of the corrosion product formed

	Elemer					
Samples	Mg	0	С	Ca	Р	Ca/P
TA TA–1PEI TA–2PEI TA–4PEI	3.57 7.02 4.59 2.67	57.54 51.73 56.94 46.67	23.56 24.77 25.39 39.11	8.01 8.94 6.96 6.55	7.11 7.55 6.11 5.00	1.13 1.19 1.14 1.31



Fig. 6 XRD curves on all corroded samples

 $(Ca_{10}(PO_4)_6(OH)_2)$  formed. Besides this, the Ca/P ratio reaches around 1.13–1.31, indicating hydroxyapatite formation.

Based on the XRD result (Fig. 6), several peaks at  $20-35^{\circ}$  and  $55-70^{\circ}$  on all samples were observed, confirming the presence of hydroxyapatite and brucite, respectively.<sup>15</sup> These results support the EDS result. Thus, this TA and TA–PEI system is suitable for the human physiological system to the evidence of hydroxyapatite formation. Moreover, the corrosion product quantification in Table 4 shows a considerable difference between the corrosion product percentage of TA and TA–4PEI, especially for hydroxyapatites. The hydroxyapatite of TA is

Corrosion product percentage (%)			
$Ca_{10}(PO_4)_6(OH)_2$	$Mg(OH)_2$		
55	45		
68	32		
40	60		
48	52		
	Corrosion product percent     Ca10(PO4)6(OH)2     55     68     40     48		

much higher than that of TA-4PEI, indicating that a more massive corrosion reaction occurred at TA compared to TA-4PEI.

## 3.4. Electrochemical corrosion behaviour of TA and TA-PEI coated pure Mg

**3.4.1. Potentiodynamic polarization curves.** The potentiodynamic polarization curve and its values of bare pure Mg, TA, and TA-PEI coated Mg are presented in Fig. 7 and Table 5.

As shown in Fig. 7, the anodic branch of polarization curves represents magnesium dissolution, and the cathodic branch represents hydrogen evolution. Pseudo-passivation in the anodic branch indicates protective layer formation and enhances the material's corrosion resistance.<sup>61</sup> Table 5 shows the data extracted from Fig. 7. The addition of TA and PEI will increase the corrosion potential value relative to pure Mg. Only the TA-2PEI sample showed a decrease in the corrosion potential value compared to TA-1PEI, and the values are relatively similar to the corrosion potential of the TA samples. It should be noted that due to the negative difference effect (NDE) in magnesium, the cathodic polarization technique is a valid method to determine magnesium's quantitative corrosion parameters instead of the anodic-cathodic approach.<sup>61-65</sup> It focused on a current density value as an essential parameter predicting the corrosion rate. Adding TA and TA-PEI coating systems onto pure Mg resulted in a declining value of the corrosion current density, which implies that the corrosion rate is decreased. The reason is that the more PEI added to the TA-coating, the coating's crack density will be lower; the coating is more



Fig. 7 Potentiodynamic polarization curve of bare pure Mg, TA, and TA–PEI coated Mg.

uniform and makes the corrosive fluid harder to penetrate further and corrode the substrate. Thus, it can be predicted that the material's corrosion resistance will be higher.

**3.4.2.** Nyquist plot and prediction of corrosion mechanisms. The EIS test results of samples that have been corroded for 1 hour, 24 hours (1 day), 72 hours (3 days), and 168 hours (7 days) in the form of a Nyquist plot are shown in Fig. 8.

As the immersion time increases, the value of impedance of all samples will increase, as shown in Fig. 8a–e, indicated by the greater semicircle in the Nyquist plot. With the increase in the impedance value, the current density that can pass through the material will get smaller. Therefore, the corrosion rate is predicted to decrease upon increasing the immersion time on the material.<sup>19</sup> The plausible cause of the increasing impedance value was the corrosion products such as magnesium hydroxide and hydroxyapatite deposited on the coating surface. These corrosion products act as a protective layer filling the crack area, hindering the SBF penetration towards the Mg surface.<sup>19</sup>

In addition, EIS characterization can also predict the evolution of different corrosion modes in the material. Based on the Nyquist plot (Fig. 8a–e), an electric circuit equivalent was made to indicate the corrosion mechanism that occurred. The electrical circuits and the corrosion mechanism in the sample are presented in Fig. 8f.

There are two models, namely model A and model B, shown in Fig. 8f.  $R_s$  is the solution's resistance between the reference and working electrode,  $C_f$  is the coating film capacitance,  $R_f$  is the resistance film,  $C_{dl}$  is the double-layer capacitance,  $R_{ct}$  is the charge transfer resistance,  $R_L$  is the inductance, and L is the resistance of inductance.<sup>19,21,28,31</sup> The primary difference between model A and model B is the presence of  $R_L$  and L in model A. These two components can be visually seen in Fig. 8a

Table 5 Corrosion potential (*E*<sub>corr</sub>), current density (*i*<sub>corr</sub>), and corrosion rate prediction obtained through potentiodynamic polarization

Sample	$E_{\rm corr}$ (V vs. $V_{\rm SCE}$ )	$-\beta_{\rm c} ({\rm mV}{ m dec}^{-1})$	$\log i_{ m corr}$ (A cm <sup>-2</sup> )	Corrosion rate (mmpy)	Ref.
Pure Mg	-2.25	0.54	-3.72	5.5	57
ГА	$-1.77\pm0.03$	$0.43 \pm 0.07$	$-3.67\pm0.05$	$4.97\pm0.06$	Experimental
FA-1PEI	$-1.60\pm0.01$	$0.39\pm0.07$	$-3.73\pm0.01$	$4.30\pm0.17$	-
FA-2PEI	$-1.76\pm0.07$	$0.24\pm0.11$	$-3.94\pm0.35$	$3.14 \pm 1.83$	
FA-4PEI	$-1.55\pm0.02$	$0.42\pm0.03$	$-3.96\pm0.29$	$2.90 \pm 1.53$	



Fig. 8 Nyquist plot obtained through EIS of samples after being immersed for 1 hour, 1 day, 3 days, and 7 days: (a) pure Mg, (b) TA, (c) TA–1PEI, (d) TA–2PEI, and (e) TA–4PEI (f) electric circuit equivalent to the Nyquist plot (top-left and top-right) and the corrosion mechanism (bottom-center).

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and b, especially in the pure Mg sample (1 hour and 1 day of immersion) and TA sample (1 hour of immersion) marked with the shape of a semicircle facing up at low frequencies. The existence of these two components indicates a localized corrosion phenomenon. Furthermore, these results align with other EIS measurements for magnesium's system as a bioresorbable implant application, thus, ensuring its applicability.<sup>19,28,29,31,40,66,67</sup>

On the other hand, the presence of  $C_{\rm f}$  and  $R_{\rm f}$  indicates the uniform corrosion phenomenon. Visually, the presence of  $C_{\rm f}$ and Rf can be seen in most of Fig. 8a-e, which is characterized by the presence of a downward-facing semicircle at high frequencies.<sup>19,21</sup> Therefore, based on EIS data and the electric circuit model equivalently, the corrosion mechanism that occurs could be predicted, as shown in Fig. 9 (bottom-center). In the pure Mg sample, the corrosion began with localized corrosion until the third day of immersion, and then uniform corrosion took place up to the seventh day of immersion. On the TA sample, localized corrosion only occurred within one hour of immersion; it continued with uniform corrosion until the seventh day of immersion. In the bare pure Mg and TA sample, the area exposed to SBF at the start of immersion was relatively high due to the absence of a coating in pure Mg and the high area of crack density in TA. As a result, localized breakdown on the sample surface caused localized corrosion to occur.<sup>19</sup> However, along with increasing immersion time (1 hour), the exposed area was reduced due to the deposition of



Fig. 9 Graphical comparison of the corrosion rate of pure Mg, TA, TA–1PEI, TA–2PEI, and TA–4PEI obtained from weight loss and EIS methods.

corrosion products on the surface so that the inductive loop (indication of localized corrosion) will slowly disappear. Thus, the corrosion mode that occurs afterward is uniform corrosion.<sup>19,21</sup> In all TA-PEI samples, uniform corrosion was the only corrosion mechanism. It is hypothesized that with a smaller crack density area in the sample at the beginning, it will be difficult for the SBF solution to penetrate into the substrate to cause a localized breakdown. In particular, when corrosion products start to be deposited, the area exposed to SBF will be reduced so that  $R_{\rm f}$  increases and  $C_{\rm dl}$  decreases.<sup>19,21,28</sup>

# 3.5. Corrosion rate measurement through weight loss and EIS methods

A comparison of corrosion rates of samples based on weight loss and EIS methods and their quantification is shown in Fig. 9 and Table 6, respectively.

As presented in Fig. 9 and Table 6, the corrosion rate prediction through the weightloss method shows the same trend. For each variation, the corrosion rate will decrease with an increase in the immersion time. If compared between samples for any given immersion time, the corrosion rate will be reduced with the increasing concentration of PEI. This is mainly due to the reduction of crack density area, which facilitates SBF penetration. In addition, the presence of corrosion product deposits on the surface also contributes to reducing the material's corrosion rate by creating a passive barrier between the substrate and SBF. Meanwhile, the corrosion rate prediction obtained through the EIS method tends to fluctuate. EIS is an instantaneous technique that predicts the corrosion rate over a certain period, i.e., when testing occurs. This method does not predict the actual corrosion rate at certain time exposure. The competition between the micro galvanic couplings formed in the substrate area exposed to SBF with the deposition of corrosion products also contributed to the fluctuating corrosion rate.<sup>19</sup>

Another concern is that the EIS method (an electrochemical method) always produces lower values of the corrosion rate compared to the weight loss method (steady state method), indicating that a non-faradaic mechanism occurred during magnesium dissolution.<sup>68</sup> This phenomenon was due to several factors. First, there is a localized reaction of formation and dissolution of MgH<sub>2</sub>. Hydrogen will cover a part of the

 Table 6
 Quantitative comparison of the corrosion rates obtained from weight loss and EIS methods

		Corrosion rate (mm year <sup>-1</sup> )					
Sample	Method	1 hour	1 day	3 days	7 days		
Pure Mg	Weight loss	18.4249	9.7306	5.5881	3.7170		
	EIS	4.0068	1.9505	0.0007	0.0013		
TA	Weight loss	15.1336	9.5707	3.2501	1.8484		
	EIS	2.2438	0.0046	0.0016	0.0015		
TA-1PEI	Weight loss	15.0156	7.2287	2.7497	1.1611		
	EIS	0.0028	0.0018	0.0013	0.0008		
TA-2PEI	Weight loss	12.4530	6.8538	2.7234	0.7092		
	EIS	0.0024	0.0006	0.0004	0.0004		
TA-4PEI	Weight loss	9.9926	7.8980	1.6347	0.4781		
	EIS	0.0019	0.0009	0.0004	0.0005		

specimen and make it unable to be detected by EIS.<sup>68,69</sup> Second is the presence of an intermediate unipositive dissolution of Mg<sup>+</sup> due to local anodic and cathodic reactions. Some of these ions will spontaneously oxidise to Mg<sup>+</sup>, and the rest will react with water to form Mg<sup>2+</sup>. This reaction causes no contribution of electron loss from the Mg substrate.<sup>69,70</sup> The last factor is the undercutting of phases. A random grain orientation in pure Mg may result in potential differences. A grain fall-off mechanism on adjacent grains may occur when the corrosion process occurs on the grain. Therefore, the grains enter the fluid physically, not as a result of an electrochemical reaction. It contributes to mass reduction but is not considered in the electrochemical methods.<sup>19,71</sup> It should be remembered that corrosion rate prediction using the Stern equation by inputting EIS data might vary depending on the chosen electrical circuit.<sup>68</sup>

Based on the explanation, the corrosion rate measured through the weight loss method will be more accurate in estimating the corrosion rate of a system. In relation to implant application, the allowable corrosion rate is 0.2-0.5 mm year<sup>-1.72</sup> Graphically, the comparison of the corrosion rate of all samples with the limitation is shown in Fig. 10.

The sample whose corrosion rate is still in the allowable corrosion rate as the bone implant is the one with a weight ratio of TA: PEI of 1:2 or coated with TA-4PEI (0.48 mm year<sup>-1</sup>) after being immersed for 7 days, as shown in Fig. 10. The reason was due to the minimum crack density area to minimize the penetration of SBF into the pure Mg substrate.

#### 3.6. Biocorrosion mechanisms

Fig. 11 shows the proposed biocorrosion mechanism at TA and TA-PEI coated pure Mg based on the actual morphology and corrosion product formed. The corrosion reaction occurred in several steps:<sup>15</sup>

(1) Through-the-thickness coating cracks will act as a medium for SBF to penetrate and react with pure magnesium as a substrate. Microgalvanic coupling between grains will be formed, and the following reaction will occur:

 $Mg(s) \rightarrow Mg^{2+}(aq) + 2e$  (anodic reaction)

$$2H_2O + 2e \rightarrow H_2(g) + OH(aq)$$
 (cathodic reaction)

$$Mg^{2+}(aq) + 2OH(aq) \rightarrow Mg(OH)_2$$
 (overall reaction)

(2) The  $Mg^{2+}$  released into the fluid will react with  $OH^-$  and form a passive layer of  $Mg(OH)_2$ . Although this passive layer is formed and deposited on coating surfaces, the protection of this layer from the corrosive fluid is not enough due to the porous structure of the layer.

(3) Due to the existence of  $Cl^-$  in SBF, Mg(OH)<sub>2</sub> will continue to be in a looping reaction of precipitation and dissolution. The site where Mg(OH)<sub>2</sub> dissolves means less protection than the area where Mg(OH)<sub>2</sub> exists. This reaction will continuously occur until there is a saturation of ions in the SBF.

(4) The site where  $Mg(OH)_2$  exists will act as a nucleation site of hydroxyapatite. With the increase of pH value and the saturation of the solution, calcium and phosphate ions will be deposited on the surface of  $Mg(OH)_2$  and form hydroxyapatite. In addition, besides an electrochemical reaction, there is a possibility of grain falling off into the solution. This evidence shows that the degradation of pure magnesium in SBF is not only controlled by an electrochemical process but also a physical degradation. The magnesium degrades to the SBF in the form of an alpha-phase-single grain instead of an  $Mg^{2+}$  ion.

## 3.7. Wettability and cell adhesion of TA and TA-PEI coated pure Mg

The results of the water contact angle test on pure magnesium samples, both uncoated and coated with four coating variations, are shown in Fig. 12a and Table 7.

As shown in Fig. 12a and Table 7, adding TA coating on bare pure Mg decreased the wetting angle from  $82^{\circ}$  to  $74^{\circ}$ . Then, along with the increase of PEI concentration in the coating system, the wetting angle value will further decrease to  $24^{\circ}$ . In other words, the wettability of the sample increases. Previous research showed that surfaces with a moderately hydrophilic characteristic  $(20^{\circ}-40^{\circ})$  exhibit the best wettability for implant application. Moreover, it will promote the highest cell attachments.<sup>73</sup> TA-4PEI is the only sample that fulfills this



Fig. 10 Corrosion rate of all samples measured through the weight loss method and the allowable corrosion rate as a bone implant.

Paper







Fig. 12 (a) Water contact angle measurement result on pure Mg samples, both uncoated and coated. (b) Cell adhesion on bare pure Mg (top left), TA coated Mg (top right), and TA-4PEI coated Mg (bottom center) observed through SEM (500×).

Table 7	Wettina	angle	values	of	all	samples
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Sample	Wetting angle (°)
Pure magnesium	$82\pm3.1$
TA	$74 \pm 1.25$
TA-1PEI	$72 \pm 4.76$
TA-2PEI	$43 \pm 4.76$
TA-4PEI	$24\pm5.96$

criterion. A decrease in the wetting angle due to the addition of TA is due to the presence of hydroxyl groups found in TA, which is considered a polar group. The polar group facilitates the presence of hydrogen bonds with water. As a result, the wettability of Mg increases. Furthermore, amino groups in PEI are also hydrophilic groups, so the coating will increasingly become hydrophilic. With the increase in PEI concentration, the number of crosslinks in the TA–PEI system will increase, and so will be the number of hydrogen bonds, which resulted in a more hydrophilic coating system. The hydrophilicity of a system is predicted to promote the adhesion of cells making the system more biocompatible.

Human Wharton's jelly mesenchymal stem cell adhesion test results of samples through SEM observation are presented in Fig. 12b. It can be inferred from Fig. 12b that on the surface of the bare pure Mg sample, there are no HWJ cells attached. However, in the TA and TA–4PEI coated samples, it is estimated that there are HWJ cells attached. In comparison, the cells grown on TA–4PEI have a better attachment and spreading than the TA-coated sample. This observation showed that the cell adhesion of TA–4PEI was better than that of TA samples. In general, it can also be concluded that the TA and TA–4PEI coating systems are proven to be non-toxic towards cells.

#### 3.8. MTT assay result

The MTT assay test result is presented in Fig. 13. The MTT test was used to measure the metabolic activity of cells seeded on the samples using (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide). This test determines the activity of cellular metabolism, that is, formazan crystal formation, indicated by the colour change from yellow to purple. This phenomenon is caused by mitochondrial dehydrogenase. The optical density (OD) value is related to cell viability; the higher the OD value, the higher the cell viability.<sup>74</sup> All samples showed increased cell viability.<sup>75</sup> The TA–PEI system offers the highest cell viability, indicating that the TA–PEI system supports the human Wharton's jelly



mesenchymal stem Cell (hWJ-MSC) proliferation and does not generate non-toxic properties compared to the others. Cells will adhere to the surface of the material in order to perform some biological activity such as migration, proliferation, and differentiation.<sup>76</sup> hWJ-MSCs have attracted great interest in bone tissue engineering since they can differentiate into the osteogenic lineage.<sup>77</sup> Previous research found that osteogenic differentiation of MSCs was observed on the scaffold with polyethyleneimine.<sup>78</sup> The scaffolds provided a suitable environment for mesenchymal Stem Cell survival and osteogenesis. In addition, corrosion of magnesium that produces hydrogen evolution and alkalization has a detrimental effect on a living organism, such as cell apoptosis if the pH value is higher or lower than 7.4.<sup>28</sup> This research proves that the higher corrosion resistance of TA-4PEI minimizes the corrosion process and alkalization. Thus, it will facilitate a higher number of cells to grow in that medium and be biocompatible and suitable for bone tissue engineering applications.

### 4. Conclusion

The results showed that a TA–PEI coating on magnesium increased the corrosion resistance and biocompatibility of the material with the following details. First, the increase in the PEI content from a weight ratio of TA:PEI 1:0 to 1:2 resulted in a reduction in the area of crack density from 25.49% to 7.12%, an increase in the coating thickness from 5.7  $\mu$ m to 9.3  $\mu$ m, a decrease in the surface roughness from 5.20  $\mu$ m to 0.98  $\mu$ m, but a change in the deformation mechanism from the ductile to the brittle layer. The reason is because of the crosslink phenomenon development between TA and PEI. Second, immersion in the SBF solution reduces the area of crack density in the TA–PEI coating. Deposits of corrosion products cover the cracks in the form of magnesium hydroxide and hydroxyapatite. Hydroxyapatite formation indicates that this system is

biocompatible with the human body. Third, adding PEI to TA as a pure magnesium coating has increased the material's corrosion resistance because the coating's crack density is reduced. Then it will make the corrosive fluids harder to penetrate and corrode the materials. Changes in the weight ratio of TA: PEI 1:0 to 1:2 reduce the corrosion rate of the material from 3.7 mm year<sup>-1</sup> to 0.47 mm year<sup>-1</sup>. In pure magnesium and coatings with a weight ratio of TA:PEI 1:0 or coated with the TA sample, the corrosion mechanism is in the form of local corrosion and then changes to uniform corrosion, while for all TA-PEI, the corrosion mechanism is uniform corrosion. The corrosion rate at a weight ratio of TA: PEI 1:2 or coated with TA-4PEI (0.48 mm year<sup>-1</sup>) meets the corrosion rate criteria allowed for bone implant applications  $(0.2-0.5 \text{ mm year}^{-1})$ . Finally, increasing the PEI content from the TA: PEI weight ratio of 1:0 to 1:2 reduces the contact angle from  $74^{\circ}$  to  $24^{\circ}$ . At a 1:2 TA: PEI ratio, passage 6 HWJ cells can adhere well and have the highest absorbance level as observed from the MTT result. Both indicate that this coating system has good biocompatibility and is potentially suitable for bone tissue engineering applications.

### Conflicts of interest

There are no conflicts to declare.

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