Improvement of Wear, Pitting Corrosion Resistance and Repassivation Ability of Mg-Based Alloys Using High Pressure Cold Sprayed (HPCS) Commercially Pure-Titanium Coatings

Mohammadreza Daroonparvar 1,*, Ashish K. Kasar 2, Mohammad Umar Farooq Khan 3, Pradeep L. Menezes 2,*, Manoranjan Misra 1 and Rajeev K. Gupta 3

1 Department of Chemical and Materials Engineering, University of Nevada, Reno, NV 89501, USA; misra49@gmail.com
2 Department of Mechanical Engineering, University of Nevada, Reno, NV 89501, USA; akasar@nevada.unr.edu
3 Department of Materials Science and Engineering, College of Engineering, North Carolina State University, 911 Partners Way, Raleigh, NC 27695, USA; mk189@zips.uakron.edu (M.U.F.K); rkgupta2@ncsu.edu (R.K.G.)
4 ASB Industries Inc., Barberton, OH 44203, USA; cmkay@asbindustries.com

* Correspondence: mr.daroonparvar@yahoo.com (M.D.); pmenezes@unr.edu (P.L.M.)

Abstract: In this study, a compact cold sprayed (CS) Ti coating was deposited on Mg alloy using a high pressure cold spray (HPCS) system. The wear and corrosion behavior of the CS Ti coating was compared with that of CS Al coating and bare Mg alloy. The Ti coating yielded lower wear rate compared to Al coating and Mg alloy. Electrochemical Impedance spectroscopy (EIS) and cyclic potentiodynamic polarization (CPP) tests revealed that CS Ti coating can substantially reduce corrosion rate of AZ31B in chloride containing solutions compared to CS Al coating. Interestingly, Ti-coated Mg alloy demonstrated negative hysteresis loop, depicting repassivation of pits, in contrast to AZ31B and Al-coated AZ31B with positive hysteresis loops where corrosion potential ($E_{corr}$) > repassivation potential ($E_{rp}$); indicating irreversible growth of pits. AZ31B and Al-coated AZ31B were most susceptible to pitting corrosion, while Ti-coated Mg alloy indicated noticeable resistance to pitting in 3.5 wt % NaCl solution. In comparison to Al coating, Ti coating considerably separated the AZ31B-Mg alloy surface from the corrosive electrolyte during long term immersion test for 11 days.

Keywords: Ti coating; cyclic potentiodynamic polarization (CPP) test; hysteresis loop; wear

1. Introduction

Magnesium (Mg) and its alloys (with a low density of about 1.8 g/cm$^3$) have become a hot topic of research because of the potential engineering applications. Moreover, magnesium alloys show outstanding potential in automotive, aerospace, and electronic industries because of their high strength-to-weight ratio, high stiffness, outstanding electromagnetic shielding ability, and remarkable damping performance, etc. In recent years, Mg alloys have been receiving ascending attention as biodegradable implant materials, as well. Regrettably, the inferior wear and corrosion performances of Mg alloys severely limit their extensive applications. The most commonly employed method for improving the surface properties of a substrate is surface treatment. In this regard, various conversion coatings [1], anodization process, plasma electrolytic oxidation (PEO), physical vapor deposition (PVD), electro-less, before processes: annealing processes, electroplating and ions implantation methods and thermal spray processes have been employed to modify the surface of Mg alloys for improving their wear and corrosion resistances [2–10].

An approximate new coating technology that deserves particular attention is cold spray process (as an environmentally friendly method) which doesn’t involve toxic fumes or other harmful emissions [11,12]. Compared to the high velocity oxy-fuel (HVOF) thermal spray process which uses a combination of thermal and kinetic energies, cold spray
utilizes only kinetic (dynamic) energy to deposit the powder particles [12-15]. Likewise, the microstructural degeneration of heat-susceptible substrates such as Mg alloys which is frequently seen in the substitute thermal spray methods could be prevented by means of cold spray process [4,16-18]. In contrast to thermal spray technologies such as electric arc wire spray, plasma spray, flame spray and HVOF spray processes which partially and/or fully melt particles during the spray process; CS can avert the thermal effects including oxidation, porosity, grain growth and phase transformation during spray process [12,13,19,20].

It was reported that corrosion resistance of Mg alloys can be improved with the cold sprayed coatings (in comparison with counterpart coatings made by other techniques e.g., anodizing, E-plating, conversion coating and etc.), in 3.5 wt % NaCl solution [21]. Aluminum (with good corrosion resistance, low density, and having low standard electrode potential difference with Mg alloys) is used (as protective coating) to reduce the corrosion rate of Mg substrates [2,22-24]. Current cold sprayed (N₂ as propellant gas) Al-based coatings (as single layer) on Mg alloys lack acceptable hardness, wear resistance and are highly susceptible to localized corrosions in severe corrosive atmospheres [22,25,26]. These coatings also showed low repassivation ability [27]. In fact, the passive film has a weak propensity to repair itself (or passivate) in corrosive environment.

Compared to Al and its alloys, Ti and its alloys can be extensively used in severe corrosive environments such as offshore (salt water), aerospace, automotive, etc. This was attributed to the good mechanical properties and excellent corrosion resistance (due to the formation of a firm protective oxide film on the metal surface) [28-30]. Low standard reduction potential mismatch between coating and substrate makes Ti coating (from group 4B) as a subsequent candidate for the corrosion protection of Mg and its alloys [31,32]. In this regard, warm sprayed (WS) Ti coatings couldn’t noticeably enhance the corrosion potential and lower the corrosion current density of AZ91E Mg alloy [31]. These coatings disclosed poor corrosion resistance and finally led to the fast degeneration of Mg alloy. The poor performance of WS Ti coatings was due to the presence of through-thickness porosities which simply conducted the chloride containing solution towards the substrate surface. The untimely tear of titanium coatings (after only 24 h of immersion) in 3.5 wt % NaCl electrolyte was eventually observed [31]. This was mainly related to the corrosion products formation and accumulation at the interface between WS Ti coating and Mg substrate [31]. The MS (magnetron sputtered) Ti-coated AZ91D Mg alloy showed even much inferior performance than the uncoated AZ91D Mg alloy after 1 day in NaCl solution [32]. Most part of MS Ti coatings came off the Mg alloy substrate which had undergone the severe corrosion [32].

In this research, we developed a fairly compact cold sprayed titanium coating on Mg alloy using HPCS system. It is anticipated that high pressure cold sprayed commercially pure-Ti coating could alleviate the problems associated with current cold sprayed Al coatings on Mg alloys and exceptionally increase the repassivation ability of Mg alloys. Moreover, immersion test for 11 days was performed to further elucidate the effectiveness and corrosion protection performance of HPCS titanium coating on magnesium alloys in corrosive environment.

2. Experimental Methods

2.1. Feedstock Powders and Substrate

In this research, commercially pure (CP) Al, and CP-Ti grade 1 powders (as feedstock powders) were employed for coating production. Commercially available AZ31B Mg alloy plate (381 mm × 455 mm × 9.5 mm) was procured from Magnesium Elektron North America. The substrates were then cut from this plate. Table 1 shows the chemical composition of AZ31B Mg alloy. The substrate surface was grit blasted and then cleaned with alcohol and acetone using a hand spray bottle and N₂ blow off right before cold spray process.
Table 1. Chemical composition of AZ31B (wt %).

| Alloy Name | Mg  | Al  | Zn  | Si  | Mn  | Fe  | Cu  | Ca  | Ni  | C  |
|------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|----|
| AZ31B Mg alloy | Balance | 2.8 | 1.1 | <0.08 | 0.7 | <0.01 | <0.01 | <0.03 | <0.001 | – |

2.2. CS Deposition

In this research, high pressure cold spray system (Impact Innovation 5/11 system, GmbH,) was used to produce metallic layers on AZ31BMg substrates. The temperature of substrate and coatings was maintained less than 65 °C during cold spray process. Likewise, Table 2 displays the cold spray parameters. The following coatings were sprayed on the AZ31B Mg alloy:

1. CS Al coating on AZ31B Mg.
2. CS Ti coating on AZ31B Mg.

Table 2. CS process parameters.

| Propellant Gas | Sprayed Material | Gas Temperature (°C) | Gas Pressure (MPa) | Spray Angle (°) | Stand-Off Distance (mm) | Step Size (mm) | Powder Feed Rate (RPM) | Powder Carrier Gas Flow Rate (m³/hr) | Type of Nozzle |
|----------------|------------------|----------------------|-------------------|----------------|------------------------|----------------|------------------------|-------------------------------------|---------------|
| N₂             | CP-Al            | 350–600              | 3.0–5.0           | 90°            | 25.4                   | 0.508          | 2.0–3.5                | 3.0                                  | SiC water cooled nozzle |
| N₂             | CP-Ti            | 700–1050             | 3.0–5.0           | 90°            | 25.4                   | 1.016          | 2.0–3.5                | 3.0                                  | SiC water cooled nozzle |

2.3. Characterization

Optical microscopy (IX70, Olympus) was used to analyze the polished cross-sectional microstructures of as-sprayed coatings on the AZ31B. For this purpose, coated AZ31B samples were cut, mounted and polished using with standard metallographic procedures on an Allied Metprep 3 TM grinder/polisher system. ImageJ software was utilized to analyze the porosity level of the as-sprayed coatings (using ASTM E2109-01) [33]. The surface morphology, polished cross-sectional microstructures and chemical composition (elemental analysis) of developed coatings before and after immersion test were studied using a LYRA-3 model XUM integrated variable pressure focused ion beam-field emission scanning electron microscope (FIB- FESEM; TESCAN), and a TM-3030 Scanning Electron Microscope (SEM; Hitachi,) equipped with Energy Dispersive X-ray spectroscopy (EDS) capability. Moreover, the structural phases of as-sprayed coatings, bare AZ31B Mg alloy substrate and feed stock powders were analyzed using an Ultima IV X-ray machine (Rigaku), after grinding up to 1200 grit size sandpaper (for only bare and coated Mg alloys). The X-ray tube emits Cu-Kα radiation with an excitation voltage of 40 kV and excitation current of 35 mA. The samples were scanned at a rate of 1 degree/min with a step width of 0.04 degree. The data were analyzed using X’Pert HighScore Plus software with ICDD database. Furthermore, 2θ (as diffraction angle) range of 30°–90° was employed to collect diffraction patterns of the different samples.

A Vickers hardness tester (Beuhler-Wilson Tukon 1202), was used to measure the microhardnesses of the substrate and the coatings, under the load of 0.245 N. It should be noted that the substrate hardness measurements were performed at the regions away from the interface between coating and the Mg alloy substrate. Additionally, ten (10) measurements were done on each sample and the average was reported as micro-hardness value.

Average surface roughness (Rₐ) of as-cold sprayed coatings were inspected during profilometry using an Alicona Infinite Focus, a 3D measurement system which has a non-contact, optical measurement principle based on focus-variation. Prior to profilometry, the surface was cleaned with DI water using an ultrasonic cleaner. The brightness and contrast were adjusted at a range of focus to make sure all the features are within focus during the scan. The lateral resolution was set at 50 nm. Rₐ was measured using line scans across the IFM scan. At least five readings were collected for Rₐ to minimize standard deviation.

Dry reciprocating sliding tests (according to ASTM G133-05 [34]) were performed using a Rtech-Tribometer at room temperature (~25 °C and 40–50% relative humidity).
Prior to sliding tests, all the coated surfaces were polished to achieve an average surface roughness (Ra) of 0.2 ± 0.05 µm. Ra is defined as the arithmetic mean of the absolute values of the vertical deviation from the mean line through the profile [35]. E52100 steel ball with 6.35 mm diameter was used as a counterpart. All the reciprocating sliding tests were carried out with a track length of 15 mm and 1 mm/s velocity under a normal load of 4 N for a total distance of 1000 mm. The 1000 mm of sliding distance was selected based on the stabilized wear depth during sliding. The wear depth was recorded to measure the wear volume. The specific wear rate was then measured using the following formula [36]:

\[
\text{Wear rate (µm}^3/\text{Nm}) = \frac{\text{Wear volume (µm}^3)}{\text{Normal load (N)} \times \text{Sliding distance (m)}}
\]

2.4. Sample Preparation for Corrosion Tests

The surface area of as-cold sprayed coatings which have a highly active surface [37] is increased. This is attributed to the rough surface of as-cold sprayed coatings. As a general practice, the rough and porous surface layer of as-cold sprayed coatings should be removed before corrosion tests [28]. Hence, the samples surface was ground up to 1200 US grit size sandpaper (SiC abrasive papers) and then cleaned with ethanol using an ultrasonic cleaner for 5 min before cyclic potentiodynamic polarization (CPP), electrochemical impedance spectroscopy (EIS) and long term immersion tests.

2.5. Cyclic Potentiodynamic Polarization (CPP) Tests in 3.5 wt % NaCl Solution

Cyclic potentiodynamic polarization tests were performed in a three-electrode setup using a flat cell and a Bio-logic potentiostat (per ASTM standard G61 [38]). The standard calomel electrode, platinum electrode, and sample under test were connected as a reference electrode, counter electrode, and working electrode, respectively. Before the CPP test, the open circuit potential (OCP) was tracked for 1 h to allow the system to achieve an equilibrium in the electrolyte. PDP tests were done in 3.5 wt % NaCl (pH 6.7) at a scan rate of 1 mV/s from 200 mV below OCP to a current limit of 10 mA/cm² or a potential limit of 2.5 V_SCE and reversed back to the same starting potential of 200 mV below OCP at room temperature. The pitting potential was determined by intersecting the line coming from extending the passive current density region and the linear increase in the current density region after passivation region.

2.6. Electrochemical Impedance Spectroscopy (EIS) in 3.5 wt % NaCl Solution

A three-electrode setup was used in a flat cell where standard calomel electrode, platinum electrode, and sample were connected as a reference electrode, counter electrode, and working electrode, respectively. The sample was monitored for 1 h observing the OCP in 3.5% NaCl (pH 6.7) exposing an area of 1 cm² at room temperature. Likewise, 100 kHz to 10 mHz (as frequency range) at OCP was selected for performing EIS test. For each EIS scan, ten measurements were recorded per decade, with an average of at least three points per measurement. Furthermore, sinusoidal AC perturbation with an amplitude of 10 mV (rms) was considered for all EIS tests. EC-lab 11.21 provided in the Bio-logic potentiostat was utilized to analyze CPP as well as the EIS data. Electrochemical corrosion tests were carried out three times to substantiate the repeatability of the obtained results. After the immersion test for 11 days, the samples were rinsed with DI water and subsequently dried in air.

3. Results and Discussion
3.1. Powders Morphology and Coatings Microstructure

CP (commercially pure)-Al (in the size range of 9–40 µm), and CP-Ti (in the size range of 10–45 µm) powders all possess spherical morphology as shown in Figure 1a,b respectively. Surface morphology of CS Ti and Al coatings (in as-sprayed condition) is shown in Figure 2b,d, respectively. Ra of coatings (in as-sprayed condition) was 2.816 ± 0.7 µm and
2.182 \pm 0.7 \mu m for Al and Ti coatings, respectively. Lower R_s for Ti coating was due to intense plastic deformation of Ti particles during CS process. Figures 2a,c and 3 demonstrate the microstructure of polished cross section of the coatings on the AZ31B substrate.

The local deformation of Ti powder particles was obvious (Figure 3c–e). Moreover, relatively dense microstructure (Figure 3a,b and Figure 2a) along with very limited micro pores with porosity level of about 0.40 \pm 0.20% was observed for titanium coating (in this research work). On the contrary, higher level of porosities were detected in atmospheric plasma sprayed (APS) Ti coating (with 10.2% porosity level) and CS Ti coating (with 2.7% porosity level) as well [31,39]. Different thermal spray methods have been employed to deposit Ti coatings with low level of porosities and high purity as well. However, thermal degradation of the deposited Ti powder particle occurs during HVOF spray process. This could be related to the temperature range of spray powder particles which is about 1227–2427 °C. In fact, the probability of a hard and brittle oxygen enriched layer formation (in the case of Ti) which is also known as “\( \alpha \)-case” is expected and could cause the notable loss of plasticity, ductility, etc. [31]. Another method is LPPS (low pressure plasma spray) process that could restrict oxidation during spray process. This is attributed to the vacuum environment which makes this technique costly. Instead, in warm spray (as modification of HVOF spray system [31]) method, supersonic gas flow temperature is adjusted by injecting N_2 into the mixing chamber. Formation of a relatively dense coating with limited oxidation of powder particles is anticipated using this method. Nevertheless, higher level of porosities was reported in the deposited Ti coatings (with porosity level of about 3.8–5.5%) on Mg alloys by WS method under different N_2 flow rates [31,39]. Higher level of porosities in warm sprayed Ti coatings substantially declined their corrosion resistance in 3.5 wt % NaCl solution and caused quick degradation of Mg substrate during long term corrosion (after 24h of immersion) [31].
Figure 2. SEM images from (a) cross section of Ti-coated AZ31B, (b) Ti coating surface, (c) cross section of Al-coated AZ31B, (d) Al coating surface, EDS analysis of 1 (e), 2 (f), and EDS analysis of 3 (g), 4 (h).

Figure 3. Cont.
Figure 3. Photomicrographs from polished crossed section of (a–e) Ti-coated AZ31B at different magnifications, and (f–j) Al-coated AZ31B at different magnifications before and after etching.

In this study, CP-Al coating with porosity level of about 1.00 ± 0.20% depicted higher porosity level than as-sprayed Ti coatings (Figure 3f–j). The presence of considerable number of micro-pores and even (worse) micro-cracks in cold sprayed CP-Al coating microstructure (at inter-particle boundaries) on AZ91D Mg alloy was also reported by Y. Tao, et al. [22]. In fact, lower degree of localized plastic deformation (localized heating, stresses) [22] resulted in an extensive formation of micro-defects at inter-particle boundaries. Cold sprayed Al coating with high denseness and having sub-micron sized grains considerably improved the corrosion resistance compared to CP-Al bulk substrates [22,25].

Figure 4 shows the XRD spectra of bare Mg alloy, feedstock powders and as-cold sprayed coatings. XRD spectrum (Figure 4a) shows that bare AZ31B is mostly comprised of α-Mg phase. Powder particles and CS coatings displayed similar phase structure and crystal planes (Figure 4b,c). Phase transformation and oxidation weren’t evidently observed in the CS coatings (Figure 4b,c). The broadened peaks in the XRD pattern of Ti (Figure 4c) coating are primarily related to the intense plastic deformation of powder particles in the coatings compared to Al coating (Figure 4b) during cold spray process [40,41]. Low processing temperature and sizable peening effect of the powder particles (during cold spray process) could lead to the retention of primary phase and crystal planes of powder particles in CS coatings. On the contrary, wire flamed sprayed Ti coatings were mainly constituted by oxides, nitrides, and carbides phases due to the nature of the flame spray process. This resulted in the inferior corrosion protection performance of the sprayed Ti coatings. Hence, these coatings had to be sealed with epoxy or Si resin for usage in the chloride containing solutions [31], so it is anticipated that a Ti coating (in this research) without any post-spray treatments could significantly decrease the corrosion rate of magnesium alloy and make AZ31B Mg alloy usable in chloride containing solutions for long periods of time.

Titanium coating considerably raised average micro-hardness (HV$_{0.025}$) of AZ31B Mg alloy surface (Figure 5a), while aluminum coating lowered average micro-hardness (HV$_{0.025}$) of substrate surface (Figure 5a). Higher micro-hardness in Ti coating may implies severe plastic deformation (high dislocation density) mostly at exterior region of powder particles (or inter-particle boundaries) that caused the increase in the coating denseness [13,25].
Figure 4. XRD patterns of (a) bare AZ31B, (b) as-received Al powder particles and as-sprayed Al coating, and (c) as-received Ti powder particles and as-sprayed Ti coating.

Figure 5. (a) Average micro-hardness (HV0.025); and (b) wear rate of bare and coated Mg substrates.

The wear depth on Ti-coated Mg alloy was around 7 μm in which Ti surface was in contact with the steel ball whereas the Al-coated Mg alloy displayed the highest wear depth of ~70 μm. The wear rate of the entire track calculated from the wear depth was plotted in Figure 5b. These results were also compared with bare AZ31BMg alloy samples that showed lower wear rate compared to Al coating surface but higher wear rate than Ti coating surfaces. In fact, a surface with higher hardness showed lower wear rate than a surface with lower hardness. This proves that CS Ti coating substantially raises surface hardness and lowers the wear rate of Mg alloys compared to CS Al coatings.
The mechanical and tribological characteristics of CS pure Ti coatings on Ti-6Al-4V substrates were studied by Khun et al. [36]. The results indicated that wear resistance of the CS Ti coating (experimented against steel balls) was noticeably higher than that of Ti-6Al-4V alloy. This was related to the cold work hardening (strain hardening) during spray process and interestingly highly wear-resistant oxide layers formation on wear tracks of CS Ti coatings (during wear tests). In fact, cold sprayed pure Ti coatings with higher compactness and lower porosities showed higher hardness and thus improved wear resistance on the Ti64 alloy as substrate [36]. Astarita, et al. also reported that CS Ti coatings can improve the wear performance of bare AA2024 alloy [42].

3.2. Electrochemical Behavior

3.2.1. Cyclic Potentiodynamic Polarization (CPP)

From the open circuit plots (Figure 6a) it can be seen that the surface of the samples was quite stable over the period of 1 h. The OCP values increase in the order: AZ31B < Al-coated AZ31B < Ti-coated AZ31B. It is evidently seen that bare Mg alloys with lower values of OCP are more susceptible to corrosion compared to coated Mg samples which showed higher values of OCP. But, Al-coated AZ31B is more active than Ti coated AZ31B.

Cyclic potentiodynamic polarization tests (Figure 6b–d) were carried out to determine whether coated and uncoated AZ31B alloys experience pitting corrosion in chloride containing solutions. This test can also help whether the passive film formed on the surface...
has a tendency to heal itself (or repassivate) in the harsh environment. Moreover, CPP test was carried out three times to prove the repeatability of the obtained outcomes.

The corrosion parameters for the bare AZ31B samples as well as Al- and Ti-coated samples are given in Table 3. It is clear that Al- and Ti-coated AZ31B alloy samples have lower corrosion current densities \( (i_{corr}) \) compared to bare samples, where lowest \( i_{corr} \) and average corrosion rate \( (P_i = 22.85 i_{corr} \) \([43]\) were observed for Ti-coated samples. The corrosion potential values were more noble for Al-coated and further higher for Ti-coated samples. Overall, it could be suggested that both Al and Ti cold sprayed coating improved the corrosion behavior of AZ31B alloy, obviously much better in case of Ti-coated samples. In this research, HPCS Ti coating considerably lowered \( i_{corr} \) to 0.049 \( \mu A/cm^2 \) from 2.504 \( \mu A/cm^2 \) and shifted \( E_{corr} \) to more noble potential, i.e., \(-387.299 mV_{SCE} \) from \(-1453.86 mV_{SCE} \) for AZ31B Mg alloy in 3.5 wt % NaCl solution. Nonetheless, magnetron sputtered Ti coating could only lower \( i_{corr} \) to 26.60 \( \mu A/cm^2 \) from 162.70 \( \mu A/cm^2 \) and \( E_{corr} \) to \(-1525 mV_{SCE} \) from \(-1570 mV_{SCE} \) for AZ91D Mg alloy in 3.5 wt % NaCl solution \([32]\).

| Type of Sample       | \( E_{corr} \) (mV \(_{SCE}\)) | \( \beta_a \) (mV/dec) | \( \beta_c \) (mV/dec) | \( i_{corr} \) (A cm\(^{-2}\)) |
|----------------------|---------------------------------|------------------------|------------------------|-------------------------------|
| AZ31B                | -1453.86                        | 115.3                  | 67.6                   | 2.504                         |
| Al-coated AZ31B      | -804.59                         | 75.5                   | 114                    | 0.092                         |
| Ti-coated AZ31B      | -387.299                        | 159.9                  | 99.7                   | 0.049                         |

Table 3. The \( E_{corr} \), \( i_{corr} \), \( \beta_a \) and \( \beta_c \) for bare and coated samples calculated from the forward scan of cyclic potentiodynamic polarization (CPP) curves.

The difference between pitting potential \( E_{pit} \) and \( E_{corr} \) (\( E_{pit} - E_{corr} \)) can be used as a measure of the propensity to the pitting nucleation \([22]\). Moreover, the difference between repassivation or protection potential \( E_{rp} \) and corrosion potential \( E_{corr} \) (\( E_{rp} - E_{corr} \)) can be employed as a measure of the repassivation ability. Larger values of \( (E_{pit} - E_{corr}) \) and \( (E_{rp} - E_{corr}) \) signify enhanced resistance to pitting corrosion and higher repassivation ability, respectively \([22]\). The \( E_{rp} - E_{corr} \) values increase in the order: AZ31B < Al-coated AZ31B < Ti-coated AZ31B. On the reverse scan, AZ31B and Al-coated AZ31B showed a positive hysteresis loop, implying the further growth of pitting. It was reported that pitting corrosion can’t get further expanded if reversed anodic curve shifted to lower current densities (as negative hysteresis loops) or the forward scan to be retraced by reversed curve. On the contrary, further pitting development is anticipated if reversed anodic curve shifted to higher current densities compared to forward scan (as positive hysteresis loops) \([44]\). The pits keep growing if \( E_{corr} > E_{rp} \) and vice versa. Ti-coated Mg alloy indicates negative hysteresis loop, depicting repassivation of pits, in contrast to AZ31B and Al-coated AZ31B with positive hysteresis loops where \( E_{corr} > E_{rp} \) indicating irreversible growth of pits. As-cold sprayed Nb coatings (from group 5B) also showed negative hysteresis loops. The repassivation behavior of CS Nb coating was attributed to the stored energy in the CS coatings assisting to passivate quickly and simply \([45]\). Analysis of \( E_{pit} - E_{corr} \) values demonstrates that AZ31B and Al-coated AZ31B are most susceptible to pitting corrosion while Ti-coated Mg alloy indicates conspicuous resistance to pitting in chloride containing electrolyte.

The CPP tests also reveal that the anodic curves for AZ31B and Al-coated AZ31B had active current densities especially in case of AZ31B alloy. The cathodic current kinetics were highest for AZ31B compared to both Al-coated and Ti-coated alloys samples. The current density limit of 10 mA/cm\(^2\) was reached at much lower potentials for AZ31B and Al-coated AZ31B alloy as well. Among all Ti-coated samples showed primary passivation followed by a secondary passivation (passive-like behavior) through a passive-active-passive (passive-like behavior) transition region. Interestingly, after the reaching the upper potential limit of 2.5 V\(_{SCE}\) the reverse scan showed lower current densities suggesting that the surface is still passive and lower potential cause lower dissolution for the passivated Ti surface. This is a very good indication of highly stable passive film which did not deteriorate even at such potentials.
3.2.2. Electrochemical Impedance Spectroscopy (EIS)

For further elucidation of the influence of cold sprayed coatings on the corrosion behavior of AZ31B Mg alloy, electrochemical impedance spectroscopy (EIS) of coated and uncoated Mg alloys was measured at their OCP. Nyquist plots are shown in Figure 7a,b. The impedance spectra was fitted by the electrical equivalent circuits (shown in Figure 7c,d: models 1 and 2). For the Nyquist diagram of uncoated sample (Figure 7d), one inductive loop and one capacitive loop at low frequency and high frequency were considered, respectively. This assumption is in harmony with the previous researches [26,46]. EEC (electrical equivalent circuit) for uncoated Mg alloy is comprised of $R_{ct}$ (charge transfer resistance), $R_s$ (electrolyte resistance), $C_{dl}$ (CPE related to electrical double layer), adsorption inductance ($L$) and adsorption resistance ($R_L$) elements.

Impedance spectra of the coated samples were fitted by using the electrical equivalent circuit, as shown schematically in Figure 8. This schematically EEC includes $R_s$ which is ohmic solution resistance at the working electrode/reference electrode interface; loop $R_{ox}$-$C_{ox}$ which shows the resistance ($R_c$ or $R_{ox}$) and capacitance ($C_c$ or $C_{ox}$) of the oxide film; $R_{ct}$ and $C_{dl}$; $R_{po}$ that is the electrolyte resistance (as additional resistance) in the localized corrosion sites (and/or the pores). The $R_c$ or $R_{ox}$ values are pretty high and any conduction of electrons through the oxide layer has been reported to be impossible [47,48]. Thus, Mansfeld and Kendig [49] proposed the removal of this circuit element from EEC and replacement of EEC1 with EEC2 [47,48]. This simplified EEC2 has been also reported in the previous researches [49–52], so a capacitive loop at high frequency and another capacitive loop at low frequency regions were assumed for the coated Mg samples. Likewise, with regard to the non-ideality of the systems, the capacitors were replaced with the constant phase elements (CPE) for all samples [51,53].

**Figure 7.** Nyquist plots of (a,b) coated and uncoated AZ31B Mg alloy substrates at OCP, electrical equivalent circuits to fit the impedance spectra of (c) coated Mg alloys (model 1), and (d) bare Mg alloys (model 2).
Figure 8. Schematic illustration of the developed EEC for fitting impedance spectra of CS Ti and CS Al-coated samples (per previous studies).

$R_{ct}$ can predominantly control the electrochemical processes rate at the interface between electrode and electrolyte (or across the electrical double layer) [54,55]. As a matter of fact, corrosion rate is reversely proportional to the $R_{ct}$ [56,57]. As presented in Table 4, Bare AZ31B Mg alloy with the lowest $R_{ct}$ showed the maximum corrosion rate between 3 samples. This could be attributed to the low protective performance of the formed corrosion surface film on the Mg alloy surface in corrosive solution. Nevertheless, values of $R_{ct}$ for Al- and Ti-coated samples were 164.707 $\Omega \cdot cm^2$, and 5580 $\Omega \cdot cm^2$, respectively. This indicates that Mg alloy could be protected by Al and Ti coatings. Nevertheless, $R_{ct}$ value for the Ti-coated Mg alloy is roughly 34 times the value of $R_{ct}$ for the Al-coated Mg alloy. The above-mentioned results reveal that the CS titanium coating can substantially declines the corrosion rate ($1/R_{ct}$) of Mg alloy in chloride containing solutions.

Table 4. EIS fitted results for bare and coated samples in 3.5 wt % NaCl electrolyte.

| Samples            | $R_s$ (\(\Omega \cdot cm^2\)) | $R_{ct}$ (\(\Omega \cdot cm^2\)) | Q_{dl} (F \cdot cm^{-2} \cdot s^{-1}) | n   | $R_{po}$ (\(\Omega \cdot cm^2\)) | Q_{c} (F \cdot cm^{-2} \cdot s^{-1}) | n   | $R_L$ (\(\Omega \cdot cm^2\)) | $L$ (H \cdot cm^2) |
|--------------------|---------------------------------|-----------------------------------|---------------------------------------|-----|---------------------------------|--------------------------------------|-----|-------------------------------|-------------------|
| Bare AZ31B Mg alloy| 18.31                           | 385.6                             | 1.13 \times 10^{-5}                   | 0.98| -                               | -                                    | -   | 187.9                         | 187.5             |
| Al-coated Mg alloy  | 15.18                           | 164707                            | 0.116 \times 10^{-3}                  | 0.87| 14397                           | 9.68 \times 10^{-6}                 | 0.88| -                            | -                 |
| Ti-coated Mg alloy  | 17.69                           | 5.58 \times 10^{6}                | 66.67 \times 10^{-6}                 | 0.798| 6059                           | 59.33 \times 10^{-6}                | 0.90| -                            | -                 |

It is worth mentioning that the corrosion resistance could be characterized by the parameter of polarization resistance at the corrosion potential ($R_{pol}$). Values of $R_{pol}$ for Al and Ti-coated samples were calculated using Equation (2) under EEC2 (Model 1) in Figure 7c [58–63].

$$R_{pol} = R_{po} + R_{ct}$$ (2)

AZ31B coated with Ti coating showed the much better corrosion resistance than Al-coated Mg alloy in 3.5 wt % NaCl solution. It is expected that this unprecedented development could be maintained even during longer immersion times.
3.3. Long Term Immersion Test in 3.5 wt % NaCl Solution for 11 Days

The surface morphological characteristics of ground coated and uncoated AZ31BMg alloys after immersion test (in 3.5 wt % NaCl solution for 264 h) are shown in Figures 9–12. Uneven corrosion products with some micro-cracks [64] (Figure 9a,b,f) entirely covered the surface of the bare AZ31B Mg alloy surface. Corrosion products on the corroded bare AZ31B Mg alloy surface were mostly comprised of Mg, O, Cl as well as Al, and Na elements (Figure 9c–e). The presence of Mg, O and Cl elements (Figure 9d,e) in the corrosion products was related to the possible existence of MgCl$_2$ and Mg(OH)$_2$ phases [57]. Nonetheless, the lower corrosion current density, higher $R_{ct}$, and $E_{corr}$ were all obtained for the AZ31BMg alloy when Al coating was applied on the AZ31B alloy (during short term electrochemical corrosion tests). Moreover, no significant corrosion was observed for the Al-coated AZ31B at the beginning of immersion test. However, deeper and even broad corrosion pits and also local net-cracks of corrosion products imply that the localized corrosion is worsened after 264 h of immersion (Figure 10a,b,f). In fact, the pitting density rises with uniform corrosion in the course of immersion time in chloride containing solutions. This behavior is also reported for the morphology of Al after corrosion [65,66].

Corrosion products on the corroded Al coating surface after 264 h of immersion were mainly constituted by Al, Mg and O as well as Cl and Na elements (Figure 10c–e). Mg element (from the AZ31B Mg substrate) was detected on the corroded Al coating surface. Al coating with low compactness could conduct chloride containing solution into the interior regions of the coating over immersion time. In fact, Al coating can’t separate the AZ31B alloy surface from the corrosive electrolyte during long term immersion for 11 days.

High pressure cold sprayed Ti (from group 4B) coating (with high propensity to repassivation and also high pitting corrosion resistance) considerably mitigated the drawbacks associated with CP-Al coating (in this research). The corroded Ti coating surface didn’t show any conspicuous corrosion pits and the other localized corrosions after long term immersion (Figure 11). It is interesting to note that, scratches (grinding tracks due to the coating surface preparation before the corrosion tests) are still distinguishable (Figure 11a,b,g) on the Ti coating surface even after 11 days of immersion.

Figure 11c–f show that corrosion products are primarily constituted by Ti and O elements which might correspond to existence of titanium-oxides [28] on the corroded coating surface. Likewise, Mg element (from AZ31B alloy substrate) wasn’t detected on the corroded Ti layer surface. In fact, the Ti layer can considerably separate the AZ31BMg alloy surface from the corrosive electrolyte even during long-term immersion for 11 days.

On the contrary, galvanic cell formation between Mg alloy substrate and warm sprayed Ti coatings were observed by Moronczyk et.al [31]. This led to the corrosion products formation and their pile up at the interface between WS Ti coating and Mg alloy substrate. This finally caused the sudden rupture of the WS Ti coatings after merely 24 h of immersion test in 3.5 wt % NaCl electrolyte [31]. This behavior was also seen for magnetron sputtered Ti coating on AZ91D after 24 h of immersion in chloride coating solution [32]. This indicates that HPCS titanium coating not only modifies the hardness and wear behavior of Mg alloys, but also can exceptionally enhance the corrosion resistance of Mg alloys in aggressive solutions during long run immersion in chloride containing solution.
3.3. Long Term Immersion Test in 3.5 wt % NaCl Solution for 11 Days

The surface morphological characteristics of ground coated and uncoated AZ31B Mg alloys after immersion test (in 3.5 wt % NaCl solution for 264 h) are shown in Figures 9–12. Uneven corrosion products with some micro-cracks [64] (Figure 9a,b,f) entirely covered the surface of the bare AZ31B Mg alloy surface. Corrosion products on the corroded bare AZ31B Mg alloy surface were mostly comprised of Mg, O, Cl as well as Al, and Na elements (Figure 9c–e). The presence of Mg, O and Cl elements (Figure 9d,e) in the corrosion products was related to the possible existence of MgCl₂ and Mg(OH)₂ phases [57]. Nonetheless, the lower corrosion current density, higher $R_{ct}$, and $E_{corr}$ were all obtained for the AZ31B Mg alloy when Al coating was applied on the AZ31B alloy (during short term electrochemical corrosion tests). Moreover, no significant corrosion was observed for the Al-coated AZ31B at the beginning of immersion test. However, deeper and even broader corrosion pits and also local net-cracks of corrosion products imply that the localized corrosion is worsened after 264 h of immersion (Figure 10a,b,f). In fact, the pitting density rises with uniform corrosion in the course of immersion time in chloride containing solutions. This behavior is also reported for the morphology of Al after corrosion [65,66].

Figure 9. (a,b) surface morphology of ground surface of uncoated AZ31B Mg alloy after immersion test in 3.5 wt % NaCl solution for 11 days, EDS analysis of 1 (c), 2 (d), 3 (e) and (f) stereo microscope image at 50× magnification.
Figure 9. (a,b) surface morphology of ground surface of uncoated AZ31B Mg alloy after immersion test in 3.5 wt % NaCl solution for 11 days, EDS analysis of 1 (c), 2 (d), 3 (e) and (f) stereo microscope image at 50× magnification.

Figure 10. (a,b) surface morphology of ground surface of Al-coated AZ31B after immersion test in 3.5 wt % NaCl solution for 11 days, EDS analysis of 1 (c), 2 (d), 3 (e) and (f) stereo microscope image at 50× magnification.

Corrosion products on the corroded Al coating surface after 264 h of immersion were mainly constituted by Al, Mg and O as well as Cl and Na elements (Figure 10c–e). Mg element (from the AZ31B Mg substrate) was detected on the corroded Al coating surface. Al coating with low compactness could conduct chloride containing solution into the interior regions of the coating over immersion time. In fact, Al coating can’t separate the AZ31B alloy surface from the corrosive electrolyte during long term immersion for 11 days.

High pressure cold sprayed Ti (from group 4B) coating (with high propensity to repassivation and also high pitting corrosion resistance) considerably mitigated the drawbacks associated with CP-Al coating (in this research). The corroded Ti coating surface didn’t show any conspicuous corrosion pits and the other localized corrosions after long...
It is interesting to note that, scratches (grinding tracks due to the coating surface preparation before the corrosion tests) are still distinguishable (Figure 11a,b,g) on the Ti coating surface even after 11 days of immersion.

Figure 11. (a,b) surface morphology of ground surface of Ti-coated AZ31B after immersion test in 3.5 wt % NaCl solution for 11 days, EDS analysis of 1 (c), 2 (d), 3 (e), 4 (f) and (g) stereo microscope image at 50× magnification.
4. Conclusions

CS Ti coatings improved the hardness and wear behavior of Mg alloys compared to CS Al coatings. AZ31B coated with Ti coating showed the much better corrosion resistance than Al-coated Mg alloy in 3.5 wt% NaCl solution. AZ31B and Al-coated AZ31B were most vulnerable to pitting corrosion, while Ti-coated Mg alloy indicated extraordinary resistance to pitting in chloride containing electrolyte. Compared to CS Al coating, cold sprayed commercially pure-Ti coating exceptionally increased the repassivation ability of Mg alloys. Moreover, immersion test for 11 days further elucidated the effectiveness and corrosion protection performance of HPCS Ti coatings on Mg alloys in corrosive solutions.

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