

Electrochemical Polarization Behaviour of Electroless Ni-P Deposits with Different Chromium-Free Pre-Treatment on Magnesium Alloy

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Different chromium-free conversion coatings such as phosphate-manganese-molybdate, vanadium and tannic-acid were used as pre-treatment for the electroless NiP deposition on magnesium alloy. The corrosion effects of NiP deposits with these pre-treatment were analyzed by electrochemical polarization tests. This revealed that NiP with tannic-based pre-treatment increased the corrosion resistance. SEM and EDS analysis revealed that it influenced not only the morphology, but also the phosphorus content of the NiP deposit. The marginal increase in phosphorus forms a layer of adsorbed hypophosphite anions, thereby preventing the hydration of nickel and forming a passive film. Furthermore, it smoothens (AFM) the NiP deposit was also the reason for the influence.

Keywords: Magnesium alloy, electroless Ni-P deposits, polarization, passive films.

1. INTRODUCTION

Magnesium and its alloys play an important role in many fields such as aerospace, electronics and automobile fields; owing to their unique characteristics of higher strength-to-weight ratio. They also have the advantages of high specific strength modulus and excellent anti-shock resistance. Therefore, they are inevitable for aerospace and automotive industries and also in manufacturing electrical equipment such as cellular phones, television sets, and sporting industries [1].

Because of their heat conductivity and electromagnetic shielding effectiveness they are also attractive features for information technology industry and for communication satellites [2]. However, it has poor atmospheric corrosion resistance and very reactive in the air, which means the magnesium oxide film will be corroded easily.

It is also hard to do electro-chemical treatment on them because of its high chemical activity to

aqueous solutions. Hence, it is not suitable to be used without any protective layer, especially in a humid environment. Furthermore, it is necessary to do surface treatment properly onto the surface to improve the adhesion, wear and corrosion resistance [3, 4].

In recent decade, different surface treatment techniques have been applied for the protection of magnesium alloys, which include conversion treatment, electroplating/electroless and anodic treatment. Conversion coatings are produced by chemical or electrochemical treatment of a metal surface to make a superficial layer of substrate metal oxides, chromates, phosphates, or other compounds that are chemically bonded to the surface [5]. There are different types of conversion coatings including chromate, phosphate/permanganate [6, 7], stannate-based [8], cerium-based, lanthanum and praseodymium conversion coatings [9, 10, 11], tannic-based coatings [12] and organic-based (silane) coatings [13].

Conversion coating is an important coating technique for magnesium alloys to resist the corrosion. Conversion coatings alone can not provide resistance from harsh corrosion, but can provide a good base for producing adherent organic coatings or paints or metallic coatings. Metallic coatings are needed for better surface finish (corrosion and wear) on magnesium alloy. Electroplating or electroless is one of the most cost effective and simple techniques for depositing a metallic coating to the magnesium substrate. In both cases, a metal salt in solution is reduced to its metallic form on the surface of the work piece.

Electroless deposition is much suitable method to deposit metal-metalloid alloy coatings than electroplating, involving the deposition with uniform, crack-free and good adhesion onto magnesium alloy, without applying an external electrical circuit. The most difficult process of plating magnesium is developing a suitable pre-treatment part. If the suitable pre-treatment is in place, many desired metals can be plated. Conventionally, the magnesium alloy was etched in a solution of chromium oxide plus nitric acid and soaked in HF solution to form a conversion film (MgF_2) before electroless plating. Due to high toxicity on environment, the hexavalent chromium compounds are restricted and HF also exhibits strong corrosive nature.

Despite the large amount of research in chromium-free coating being carried out for many years, there have been a few products that can provide excellent corrosion protection with low cost treatment until now. The phosphate-manganese-molybdate coating (PMMC), vanadium based coating (VBC) and tannic based coating (TBC) were used in these study. Our previous [4] study mainly emphasis on the variation in time and concentration of these conversion coatings and electroless NiP deposits has not been well established. Hence, the present research provided an opportunity to emphasis more analysis on the electroless nickel-(EN)-phosphorous (Ni-P) deposit with using these pre-treatments. As several properties of metal deposited magnesium alloy are dependent on the good pre-treatment, the main aim of this present study was to understand the effect of different pre-treatments on the corrosion behaviour of electroless NiP depositions. These chromium-free conversions coating was studied by XPS and X-RD. The microstructures and the electrochemical properties of the EN deposit with these chromium-free pre-treatment on the AZ91D Mg alloy substrate were studied by SEM, X-RD, AFM and electrochemical polarization measurement.

2. EXPERIMENTAL PROCEDURE

The substrate material used was AZ91D die cast magnesium alloy with a size of 15×15×5 mm. The alloy was mainly contained about 9.1 wt. % Al, 0.64 wt. % Zn, 0.17 wt. % Mn, 0.001 wt. % Fe and Mg balance. The substrate was ground with No. 2000 SiC paper.

Table 1. The operating parameters of alkaline cleaning on the AZ91D magnesium alloy.

Process	Parameters	Conditions
Grinding	Up to No. 2000 SiC sandpaper	
Alkaline cleaning	NaOH - 45 g/L Na ₃ PO ₄ ·12H ₂ O - 10 g/L	338 K, 20 min

After grinding, the sample was rinsed with deionized water, cleaned in acetone ultrasonically for a while and finally dried, before take into the degreasing process. After degreasing (given in Table-1), the above-mentioned chromium-free pre-treatments were applied before the electroless NiP plating. Magnesium and its alloys have a high affinity to oxygen, which means that the freshly cleaned samples will always be covered with a thin oxide film growing rapidly. So, samples should be transferred as quickly as possible between any two procedures.

Table 2. Pre-treating bath for Phosphate-manganese-molybdate pre-coating for AZ91D alloy.

Phosphate-manganese-molybdate coating (PMM)*	Conditions
Mn(H ₂ PO ₄) ₂ - 0.5 g/L H ₃ PO ₄ (85% V/V) - 15 ml/L C ₂ H ₄ O ₂ - 20 ml/L CH ₃ CH ₂ OH - 20 ml/L HNO ₃ (80 % V/V) - 5 ml/L Na ₂ MoO ₄ - 2.5 g/L	Time 3 min at 298 K

*Phosphate-manganese-molybdate coated pre-treatment sample is represented by AZ91D/PMMC

The phosphate-manganese-molybdate coating (PMMC) (given in Table-2) is a modification of the common phosphate-manganese pre-treatment procedure. Because, the addition of molybdenum from sodium molybdenum can disintegrate into MoO₄²⁻ ions, which could combines with metal ions to form MgMoO₄ as follows.



And then, this behave as the cathode sites (in similar with magnesium matrix) and stimulate the phosphates-manganese conversion coating [14]. This is similar to the case of phosphating on steel, where MoO_4^{2-} ions coupled with Fe^{2+} to form FeMoO_4 that is absorbed onto the iron substrate to behave as nuclei to stimulate the growth of phosphates [15]. Hence, the addition of molybdate could create more suitable cathode sites for the facilitating and growth of phosphate film. As a consequence, a fine and fully covered phosphate film was formed on the substrate.

Table 3. Pre-treating bath for Vanadium pre-coating for AZ91D alloy.

Vanadium based coating pre-treatment (VBC)*	Conditions
NaVO_3 - 30 g/L	Time 15 min at 353 K, pH 8.0.

*Vanadium based coating pre-treatment sample is represented by AZ91D/VBC.

Table 4. Pre-treating bath for Tannic acid based pre-coating for AZ91D alloy.

Tannic based coating pre-treatment (TBC)*	Conditions
$\text{C}_{76}\text{H}_{52}\text{O}_{64}$ - 0.8 ml/L	Time 5 min at 298 K, pH 4.00
NH_4VO_3 - 1 g/L	
K_2ZrF_6 - 1.1 g/L	
Na_3PO_4 - 1 g/L	
HNO_3 - 0.4 ml/L	
$\text{Na}_2\text{B}_7\text{O}_4$ - 5 g/L	

*Tannic based coating pre-treatment sample is represented by AZ91D/TBC.

Another pre-treatment technique applied is vanadium based coating (VBC) (given in Table-3). Vanadium conversion is generally used as the corrosion resistant inhibitor for the paint or pigment system, it is scanty to be used on magnesium alloy. The vanadium coating formation process is gelatine of hydrated vanadium oxide, which is softer than those by other methods, and easy to control. The formation of pentavalent vanadium oxides by condensation and polymerization has been responsible for the formation. Vanadium oxide gel formation includes ovation of octahedrally coordinated $\text{VO}(\text{OH})_3(\text{OH}_2)_2$. Yang et.al had developed the vanadium based conversion coating on the AZ61 magnesium alloy and reported that corrosion resistance of vanadium coating is better than cerium and phosphate coatings, and in addition this vanadium chemical conversion treatment has potential to replace the chromium-based conversion coating treatment [16]. The vanadium treatment of 10 min time was good for AZ61 alloy and 15-20 min time was good for AZ91D case. The increasing β -phases in this AZ91D case might be the reason for the delay in the coating time [17].

And finally, an unfamiliar pre-treatment called tannic based coating for AZ91D alloy (given in Table-4). The tannic acid based conversion coating can be formed on AZ91D magnesium alloy

through a solution containing $C_{76}H_{52}O_{46}$ (tannic acid), NH_4VO_3 , K_2ZrF_6 and H_3PO_4 . Pentahydroxy benzamide-magnesium complex was the main component of the coating, which is succeeding in two steps: $Zr(HPO_4)_2 \cdot H_2O$ (α -ZrP) formed first at ~70 to 120 second of immersion time, which behaved as a catalyst to activate the main reaction. The oxidization of gallic acid into pentahydroxy benzoic acid was the second reaction, as well as the formation of pentahydroxy benzamide-magnesium complex, which took place in the time interval of about 120 to 300 s. The reactions take up first on β phases and then spread to α phase [12].

Table 5. Bath composition and operating parameters of electroless NiP deposits.

Electroless NiP deposition parameters*	Conditions
NiSO ₄ ·6H ₂ O - 0.06 mol/L	pH 6±0.2 at 355±2 K.
NaH ₂ PO ₂ ·H ₂ O - 0.15 mol/L	
HF (40% V/V) - 0.6 mol/L	
NaC ₂ H ₃ O ₂ - 0.16 mol/L	
NH ₄ HF ₂ - 0.14 mol/L	
Stabilizer - 1 ppm/L	

*Electroless nickel plating samples with these different pre-treatment are represented by AZ91D/PMMC/EN, AZ91D/VBC/EN and AZ91D/TBC/EN.

After being pretreated in the above pre-treatment baths separately, the samples were immersed in the electroless bath solution for NiP deposition layer. The technical flow chart of the electroless NiP deposition is shown in Table-5. The samples were cleaned thoroughly with de-ionized water as soon as possible between any two steps of the treatments. The electroless solution was put in a 1000 ml glass beaker, which was kept at constant temperature by a thermostat. The bath composition and all the operation parameters for the electroless NiP deposition are listed in Table-5. The thickness of Ni-P deposition was calculated by weight gain method and it was found to be around 25 μ m.

The structures were studied by the X-ray diffractometer (XRD, Rigaku Dymax) with a Cu K α radiation ($\lambda=0.154178$ nm) and a monochromator at 50 kV and 300 mA with the scanning rate and step being 4°/min and 0.02°, respectively. X-ray photoemission spectroscopy (XPS) was used to analyze the component of the conversion coatings prior to electroless NiP depositions. XPS spectra of the coatings were acquired with an ESCALABMk II (Vacuum Generators) spectrometer using un-monochromatized Al K α X-rays (240 W). Cycles of XPS measurements were done at high vacuum chamber with a base pressure of 10⁻⁸Torr. The surface morphology of the electroless nickel coatings were observed by using scanning electron microscope (SEM, JEOL JSM-5310) and an EDS attachment was used for qualitative elemental chemical analysis. The surface topography of the electroless coated samples was observed by atomic force microscope (AFM) and their histogram reveals the surface roughness (R_a) value of the electroless NiP samples.

Electrochemical polarization measurements were performed on an electrochemical analyzer

(Versa STAT-3, METEK, Princeton Applied Research), which was controlled and supported by a computer software system.

Polarization (Potentiodynamic, Tafel and linear) experiments were conducted at ambient temperature ($\sim 25^{\circ}\text{C}$). The experiments were carried out in a 3.5 wt. % NaCl aqueous solution using a three-electrode cell with a platinum plate (Pt) as counter electrode and a saturated calomel electrode (SCE, +242 mV vs. SHE) was fitted through the rubber stopper and used as a reference electrode. The exposed area for testing was obtained by thick coating the sample with epoxy resin leaving an uncovered area of 1 cm². The reference and platinum electrodes were fixed near to the working electrode (~ 0.5 mm), which could minimize the errors due to IR drop in the electrolytes. All the experiments were performed with the scan rate of 5 mV/s. The log (i)–E curves were measured and plotted after the above electrochemical measurements. The free corrosion potential was stabilized prior to each potentiodynamic and anodic polarization experiment. However, the Tafel experiments were conducted immediately on immersion of the sample in the electrolyte in order to understand the corrosion resistance of fresh surfaces of EN deposits on magnesium alloy and also avoid passivation to measure the corrosion rate.

3. RESULTS AND DISCUSSION

3.1. The study of chromium-free treatments on AZ91D Mg alloy

The XRD patterns of the AZ91D magnesium alloy substrate, the three different chromium-free pre-treated surfaces are shown in Fig.1.

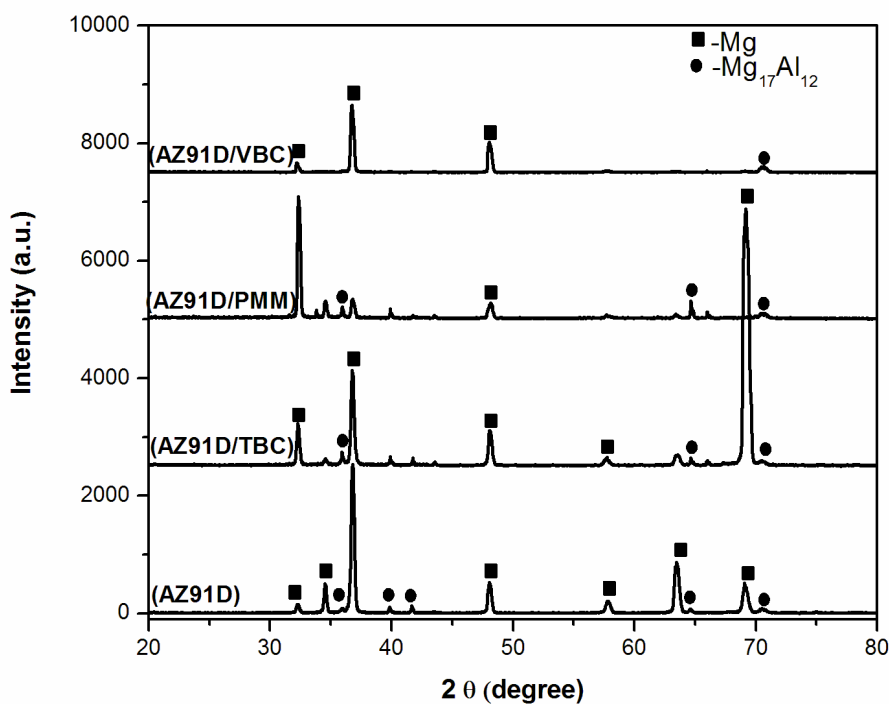
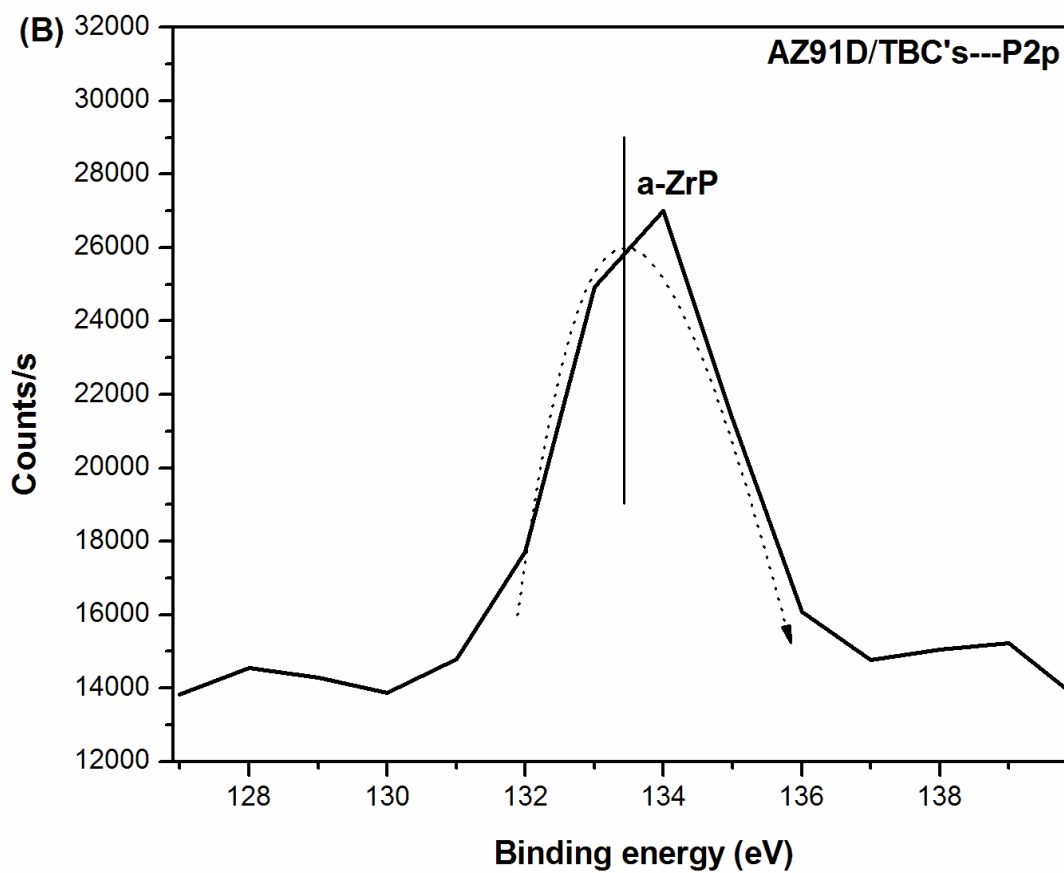
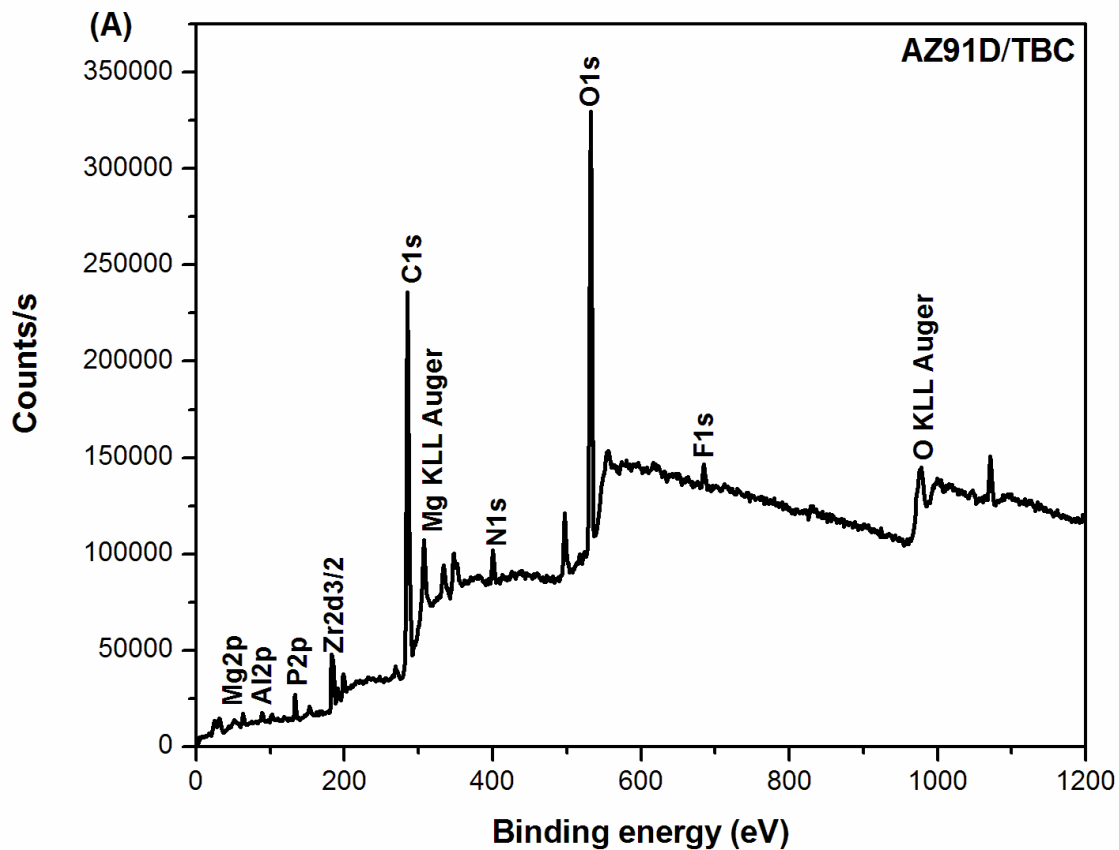


Figure 1. XRD patterns of AZ91D and three different chromium-free pre-treated on AZ91D magnesium alloy substrate.



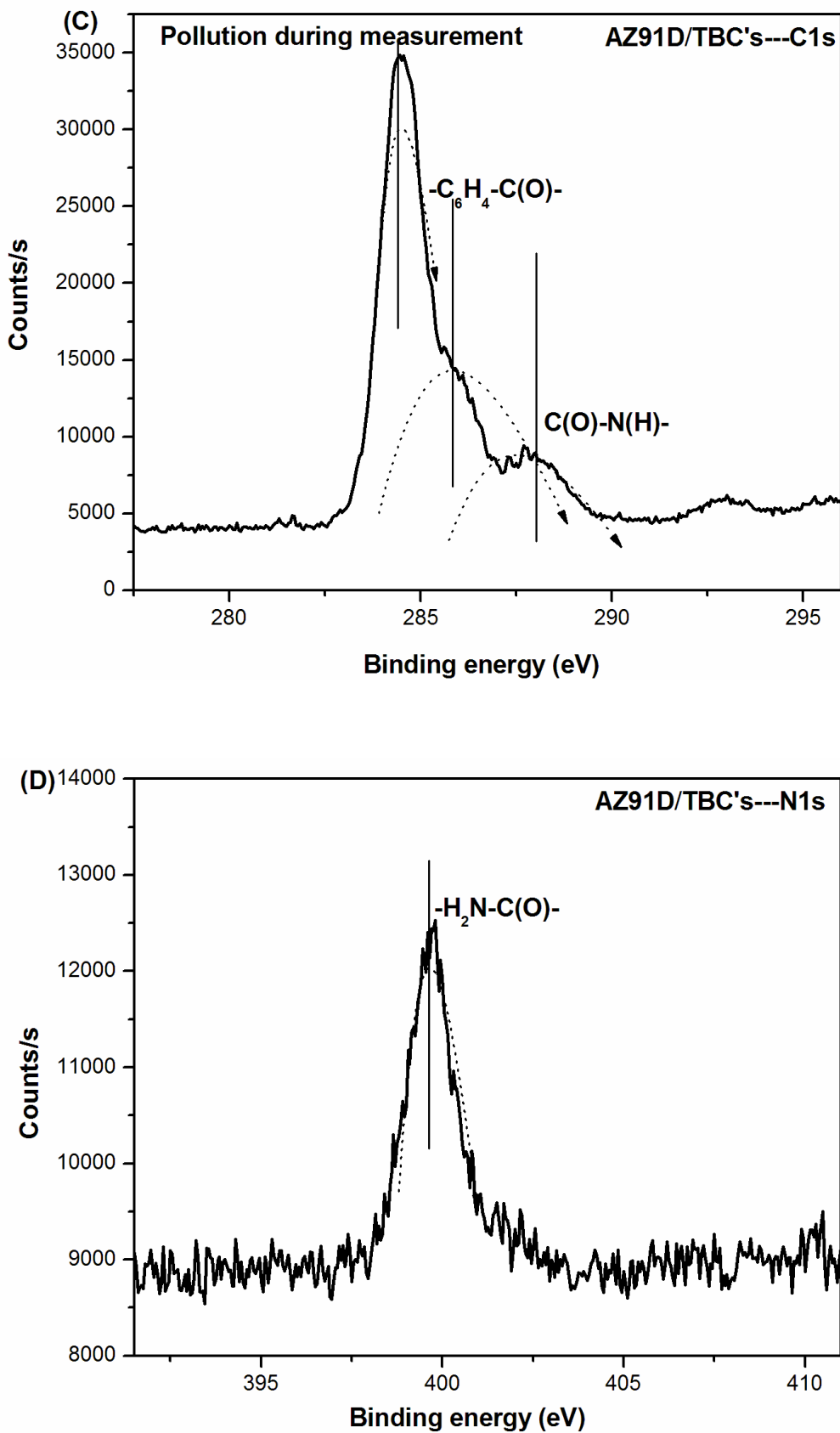


Figure 2. XPS spectrum of AZ91D magnesium alloy substrate treated in tannic based pre-coating: (A) survey spectra, (B) enlarged P2p XPS spectrum, (C) enlarged C1s XPS spectrum, (D) enlarged N1s XPS spectrum (at ambient conditions and prior to EN deposit).

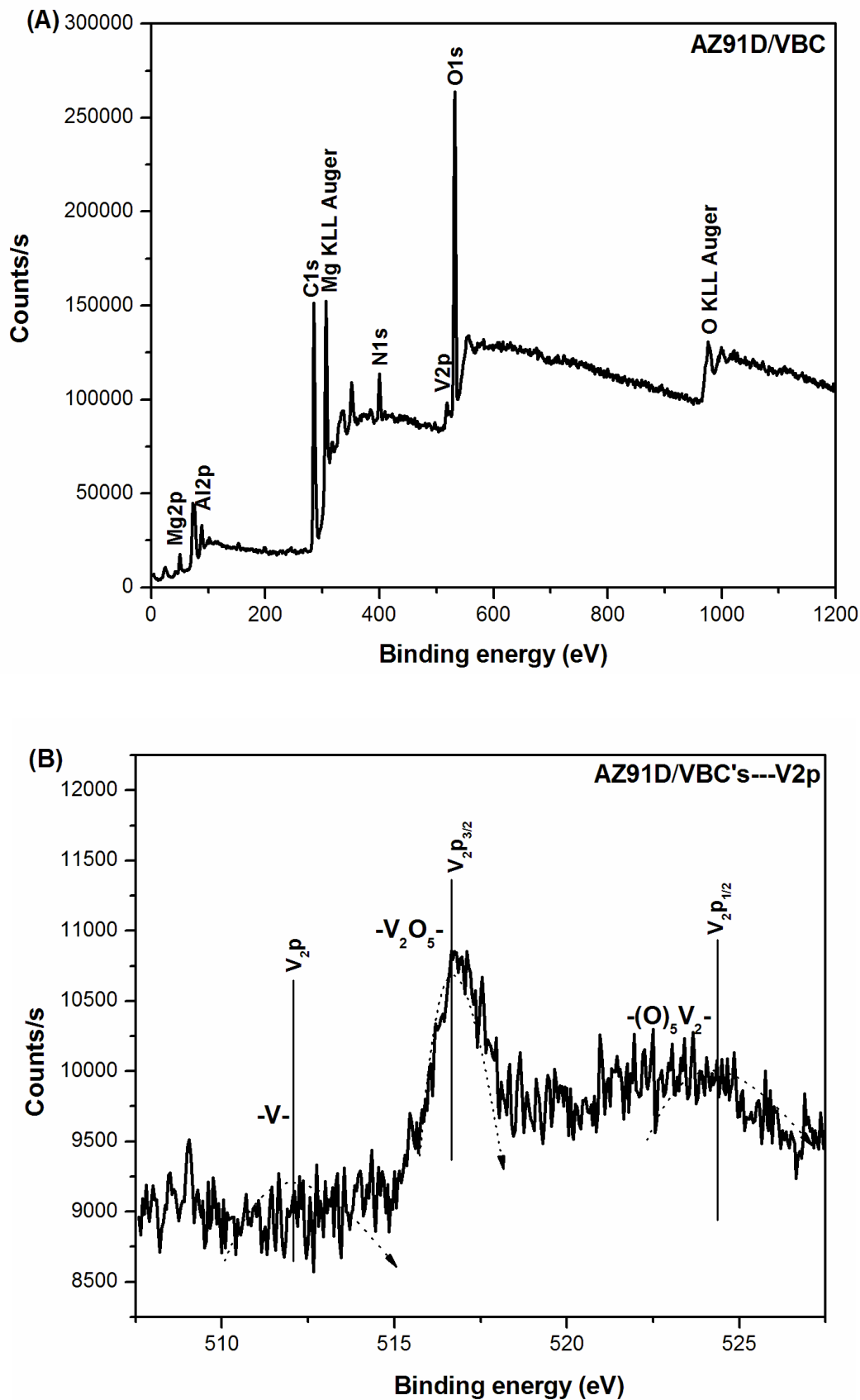


Figure 3. XPS spectrum of AZ91D magnesium alloy substrate treated in vanadium based pre-coating: (A) survey spectra, (B) enlarged V2p XPS spectrum (at ambient conditions and prior to EN deposit).

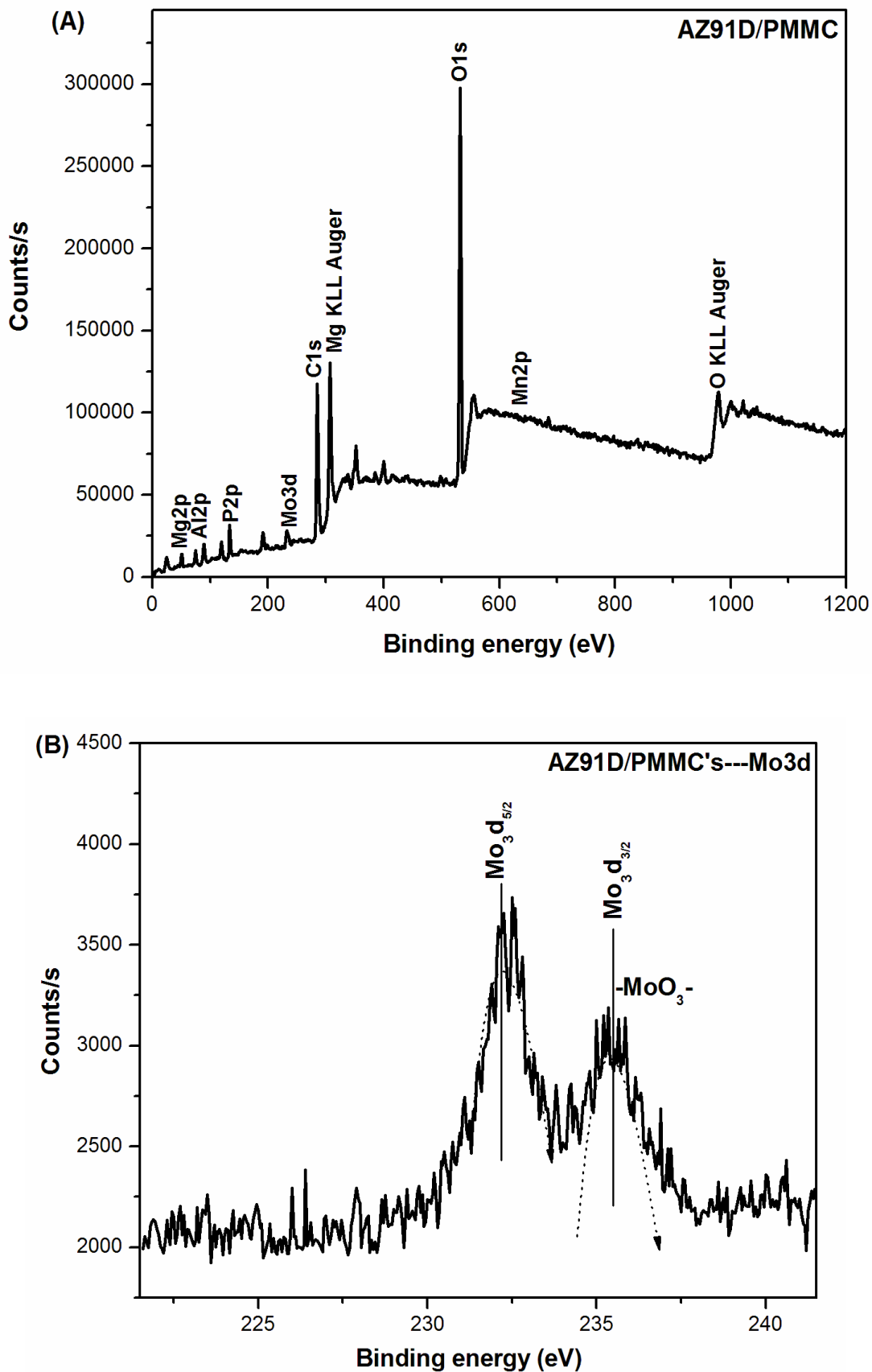


Figure 4. XPS spectrum of AZ91D magnesium alloy substrate treated in phosphate-manganese-molybdate pre-coating: (A) survey spectra, (B) enlarged Mo3d XPS spectrum (at ambient conditions and prior to EN deposit).

Conversion coating thickness is less than 1 μm , hence, the phases in the pretreatment layer detected very less by XRD; the pale colored surface may imply the existence of passive film on the substance. At the same time, the AZ91D magnesium alloy consisted of primary α -Mg phase surrounded by a eutectic mixture of α and β - $\text{Mg}_{17}\text{Al}_{12}$ [18]. There is an internal galvanic corrosion caused by the second phases or impurities. The β -phase precipitated along the grain boundaries, which exhibited higher cathodic reaction activity and lower corrosion current density than that of α [18]. Hence, a suitable pre-treatment is in need to make sure for the successful metallic coating on magnesium alloy. After pre-treatment, α phase was slightly eroded and β phase was exposed on the surface. This results in the increase of the volume fraction of β phase on the surface. Both the precipitation of conversion coatings and the increase of the volume fraction of β phase can improve the corrosion resistance of the substrate, because the galvanic corrosion is reduced [19].

The XPS was used to analyze the component of the conversion coatings prior to electroless NiP deposits. The spectra of the different chromium-free pre-treated samples were shown in Figs. 2-4, respectively. XPS spectrum of Mg2p, Al2p, C1s and O1s peaks are commonly seen in all the three pre-treated samples. Fig. 2(A) shows the XPS spectrum of tannic based coating on AZ91D substrate (AZ91D/TBC). Both the P2p peak (Fig. 2(B)) at 133.44 eV and the Zr3d5/2 peak at 182.9 eV are assigned to α -ZrP [20]. However, these both peaks will form only in the early stage of conversion coating, which may act as catalyst to facilitate the subsequent reaction of gallic acid oxidation into pentahydroxy benzoic acid. The C1s spectrum (Fig. 2(C)) reveals three peak structures with binding energies at 284.5 eV, 285.85 eV and 288.01 eV respectively. The peak at 284.5 eV is caused by pollution during XPS measurement. The peak at 285.85 eV is attributed to $-\text{C}_6\text{H}_4-\text{C}(\text{O})-$ group in the organic polymer [21], which should be the reaction product of tannic acid. The N1s peak (Fig. 2(D)) locates at 399 eV assigned to the $-\text{H}_2\text{N}-\text{C}(\text{O})-$ group [22]. The Mg2p peaks at 49.6 eV and 51.0 eV are observed, which are attributed to the Mg-Mg bond [23] and Mg-F bond [24], respectively. Fig. 2(A) also shows the peaks of O1s, Mg2p and F1s obtained on sample treated for TBC. Mg2p and F1s peaks are attributed to substrate Mg and fluoride in the conversion coating, respectively. Fig. 3(A) shows the XPS spectrum of vanadium based coating on AZ91D substrate (AZ91D/VBC). The V2p peak (Fig. 3(B)) reveals three peak structures with binding energies at 512 eV, 516.65 eV and 524.3 eV respectively. The peak at 512 eV is attributed to $-\text{V}$ -metallic vanadium bond [25]. The peak at 516.65 eV and 524.3 eV are attributed to $-\text{V}_2\text{O}_5-$ group in the metallic oxide form [26]. The Al2p peak reveals that Al_3V bonds are present in the 88.5 eV range. Fig. 4(A) shows the XPS spectrum of phosphate-manganese-molybdate coating on AZ91D substrate (AZ91D/PMMC). Apart from the Mg2p, Al2p P2p, C1s and Mn2p peaks, the Mo3d peak is important in this PMMC pre-treatment and it (Fig. 4(B)) reveals two-peak structures with binding energies at 232.9 eV and 235.3 eV. The both peaks are attributed to $-\text{MoO}_3-$ group in the metallic oxide form [26]. The Mn2p peak is also present at 642.12 eV range.

The SEM images of these conversions pre-treatment on magnesium alloy can be seen from our earlier research [4, 12, 17]. It can be seen from the above XPS analyses that the conversion treatments forming successfully on AZ91D alloy and some of them preferentially nucleated on the β phase and then extended to other phases of the substrate. Both the precipitation of conversion coatings and the

increase of the volume fraction of β phase can improve the corrosion resistance of the substrate, during electroless NiP deposit. The effect of electrochemical polarization (potentiodynamic, anodic and tafel) results on these three chromium-free pretreatment layers for electroless process is discussed in coming section.

3.2. The electrochemical properties of electroless NiP with different pre-treatments

The three different chromium-free treatments were successfully used as a pre-treatment for electroless NiP alloy coating on the AZ91D Mg alloy substrate. The XRD pattern of the electroless NiP deposit with different pre-treatment on AZ91D alloy is shown in Fig. 5.

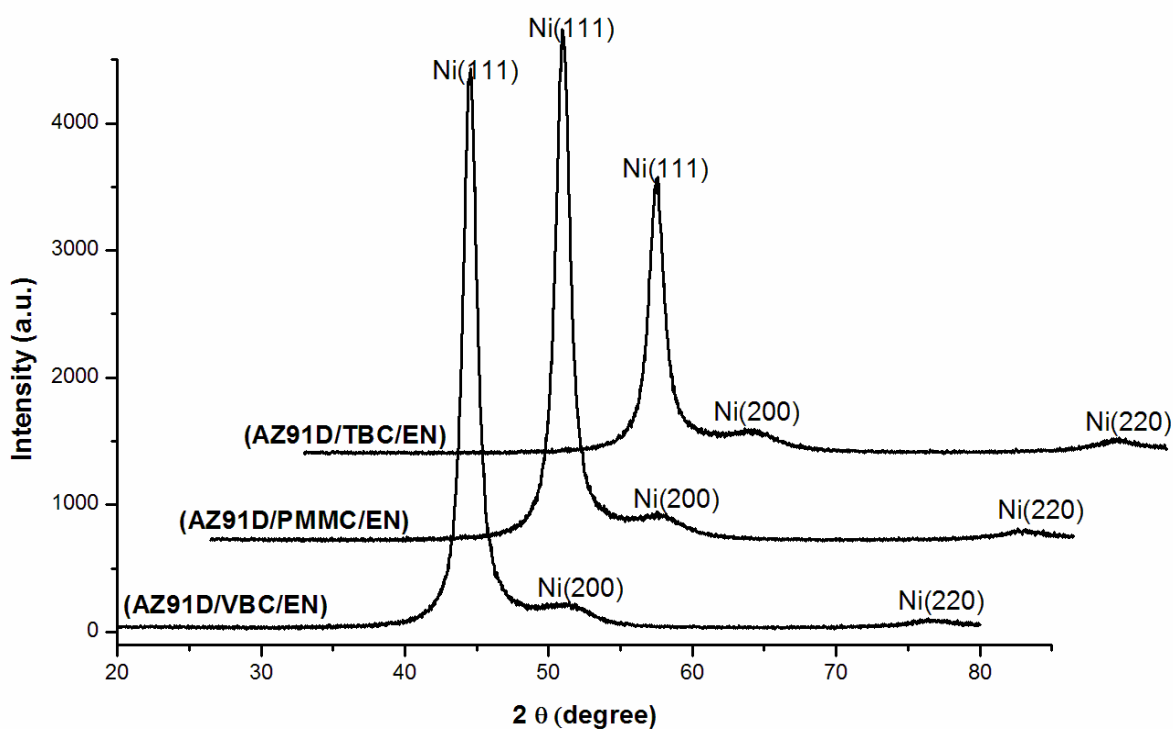


Figure 5. XRD patterns of EN deposit with three different chromium-free pre-treated on AZ91D alloy.

The EN deposits show very broad peak corresponding to the (111) plane of a face-centered cubic (fcc) phase of nickel ($2\theta=44.58^\circ$, $FWHM=0.75\sim 0.83$) and two small broad peaks of Ni(200) and Ni(220), which in fact indicate the mixture of nanocrystalline and amorphous structure. And small grain size has very great impact on corrosion performance [27]. Lo et.al had studied the role of phosphorus in the electrochemical behavior of electroless Ni-P alloys in 3.5 wt. % NaCl solutions [28]. It showed that high P content coating shows amorphous structure and improved corrosion resistance. The enriched phosphorus surface reacts with water to form a layer of adsorbed hypophosphite anions [29]. This layer will in turn block the supply of water to the electrode surface, thereby preventing the hydration of nickel, which is considered to be the first step to form either soluble nickel ion species or a passive nickel film [28], which is discussed in detail in coming section. In this study, the phosphorus

content by EDS analysis (not shown) reveals that all three samples are in the medium range (~5.8 wt. %) of the mixture of nanocrystalline and amorphous structure [30]. But there was a relatively high 'P'-content for the sample AZ91D/TBC/EN than other cases. It is generally accepted that the relative ratio of nanocrystalline and amorphous or a co-existence of these two phases depends on the phosphorus content [30].

The surface morphology of electroless NiP deposits with different pre-treatments is shown in Fig. 6 (A, B & C).

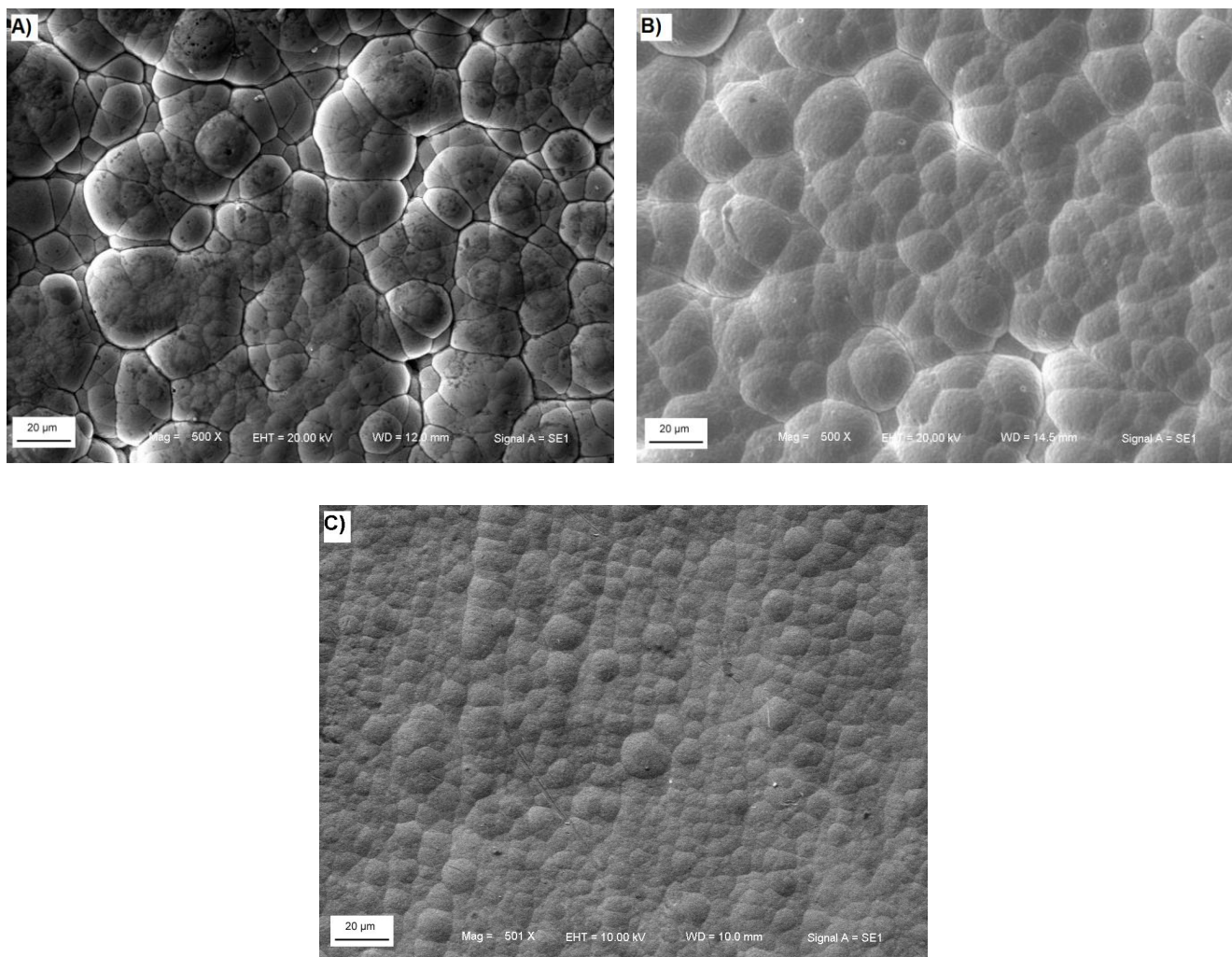


Figure 6. SEM images of EN deposit with: (A) Phosphate-manganese-molybdate coating, (B) Vanadium based coating, (C) Tannic based coating on AZ91D alloy.

The deposit surface morphology reveals that all the electroless NiP deposits show globular appearances, which are generally compact, porous-free and uniform in size. All the pre-treatments do have influence on the deposit surface morphology. It is seen from the figure-6 that the EN deposit with VBC treatment is more compact than that of PMMC, while the NiP deposit with TBC shows the smallest globular and the smoothest and compact surface among the three coatings. These facts are in accordance with that the pre-treatments should have some effects not only on the initial stage of

deposition, but also on the whole catalytic EN deposition process. Thus, it is believed that by changing the surface structure and size of active area of substrate, it can influence the nucleation rate of electroless NiP at the initial stage. In addition, the lateral joining of the growing crystallites can also be evidently influenced. Hence, not only the morphology, but also the micro porosity and other inhomogeneities (e.g. degree of internal stress and type) of the EN deposition will be influenced as well.

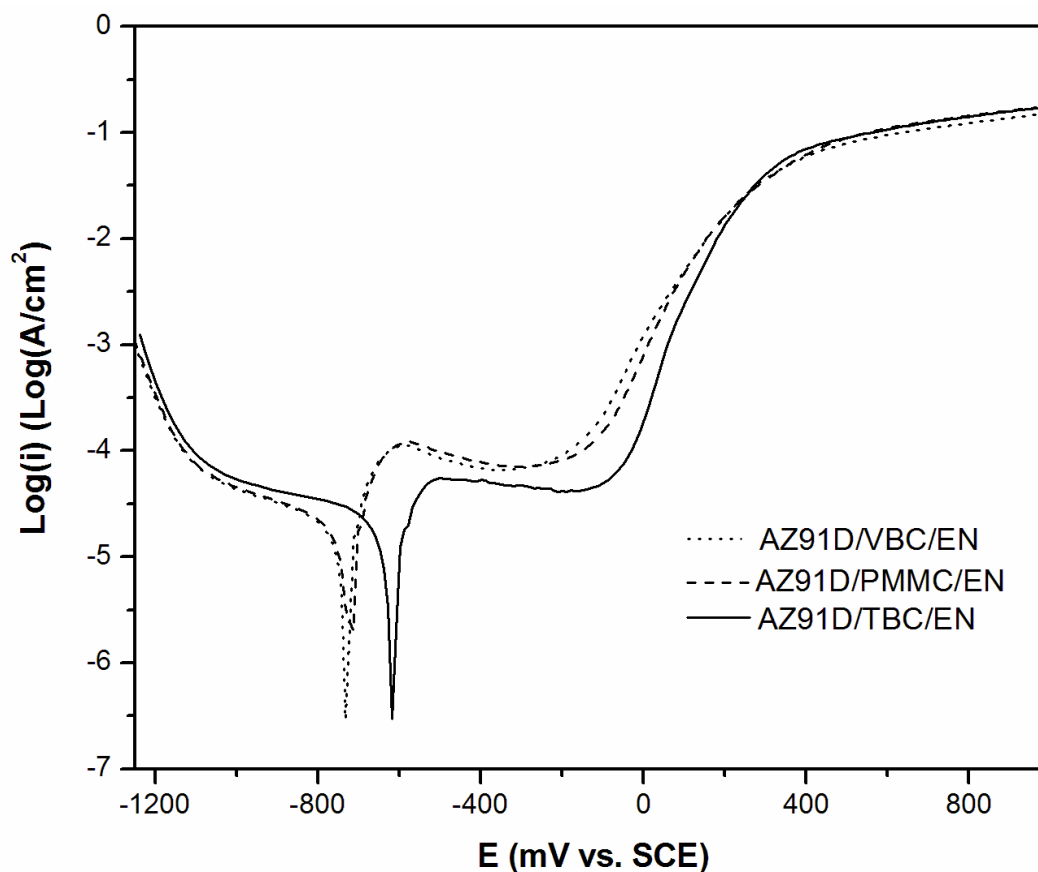


Figure 7. Potentiodynamic polarization curves of different chromium-free pre-treated EN deposit on AZ91D magnesium alloy. The experiments were conducted after stabilization of free corrosion potentials (conducted at room temperature in 3.5 wt% NaCl at a scan rate of 5 mV/s).

Fig. 7 shows the potentiodynamic polarization curves of the EN deposited magnesium alloy with different chromium-free conversion coatings in 3.5 wt. % NaCl solution at room temperature. Active-passive polarization behaviors were observed for all EN deposited magnesium alloy (Fig. 7). The cathodic process occurred in the cathodic branch, and the main reaction was hydrogen evolution. The experiments were conducted after stabilization of free corrosion potentials and they were reproducible, based on two experiments conducted for each case. As the nickel deposit was cathodic relative to the substrate, the positive shift of free corrosion potential of the alloy with these conversions coating will reduce the potential difference between the nickel/substrate. It thereby improves the corrosion resistance of the conversion coating plus EN deposition and avoids the catastrophic galvanic corrosion between nickel and magnesium. From the above electrochemical potentiodynamic

polarization curve the passivation parameters, i.e., zero current potential (ZCP), potential for primary passivation (E_{pp}), potential for complete passivation (E_{cp}), critical current density (i_{crit}), passive current density (i_{pass}), passive range ($E_b - E_{cp}$) and break-down potential (E_b) are presented in Table-6.

Table 6. Passivation parameters obtained from polarization curves of different chromium-free pre-treated EN on AZ91D alloy in 3.5 wt. % NaCl at room temperature.

Passivation parameters	AZ91D/VBC/EN	AZ91D/PMMC/EN	AZ91D/TBC/EN
ZCP (mV vs. SCE)	-731	-714	-618
E_{pp} (mV vs. SCE)	-676	-669	-577
E_{cp} (mV vs. SCE)	-583	-590	-515
E_b (mV vs. SCE)	-241	-213	-107
$E_b - E_{cp}$ (mV)	342	377	408
i_{crit} ($\mu\text{A}/\text{cm}^2$)	30.4	26.1	20.4
i_{pass} ($\mu\text{A}/\text{cm}^2$)	110.6	118.9	45.2

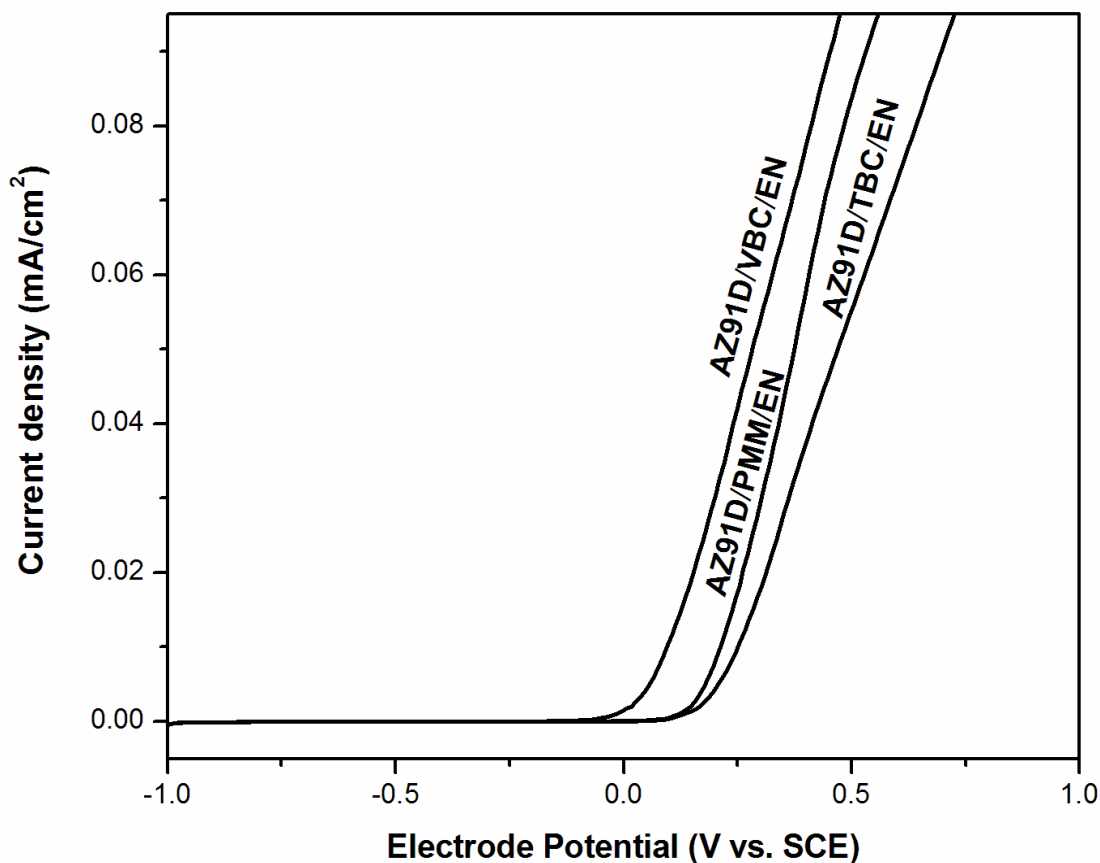


Figure 8. Anodic polarization curves of different chromium-free pre-treated EN deposits on AZ91D magnesium alloy. The experiments were conducted after stabilization of free corrosion potentials (conducted at room temperature in 3.5 wt% NaCl at a scan rate of 5 mV/s).

The critical current densities and passive current densities were determined at E_{pp} and E_{cp} , respectively. In specific, there was a shift in ZCP towards positive direction for AZ91D/TBC/EN compare to those with PMMC and VBC. The positive shift in the ZCP may be attributed to changes in hydrogen reduction processes [31], due to substantial fraction of atoms lie in the inter-crystalline region of nano crystalline nickel. The critical current densities of passivation for all the samples were approximately similar. This indicates that the mechanism of onset of passivity and passivation itself is similar in nature for all the case.

There was a noticeable change in the passive range for TBC compared to other cases. There was a significant change in the passive current density for TBC compared to PMMC and VBC. This might be non-defective nature of passive film formed on TBC. The reason may be related to the relatively high (than VBC and PMMC) phosphorus content on the surface reacts with water to form a layer of adsorbed hypophosphite anions. Hence, the passive film formed by TBC passive film has positive influence [28]. When the applied anodic potential was high enough to breakdown the passive film, pitting corrosion occur at the grain boundaries. Fig. 8 shows anodic polarization curves recorded of the EN deposited magnesium alloy with different pre-coatings in 3.5 wt. % NaCl solution at room temperature. The experiments were conducted after stabilization of free corrosion potentials. As the anodic potential increases towards positive direction, the corrosion current started to increase slowly. The current plateau is clearly associated with the metallic oxidation. The dissolution of metal matrix differs marginal for EN deposited with TBC. This behavior indicates similar like that of the marginal increase in the potentiodynamic polarization test also.

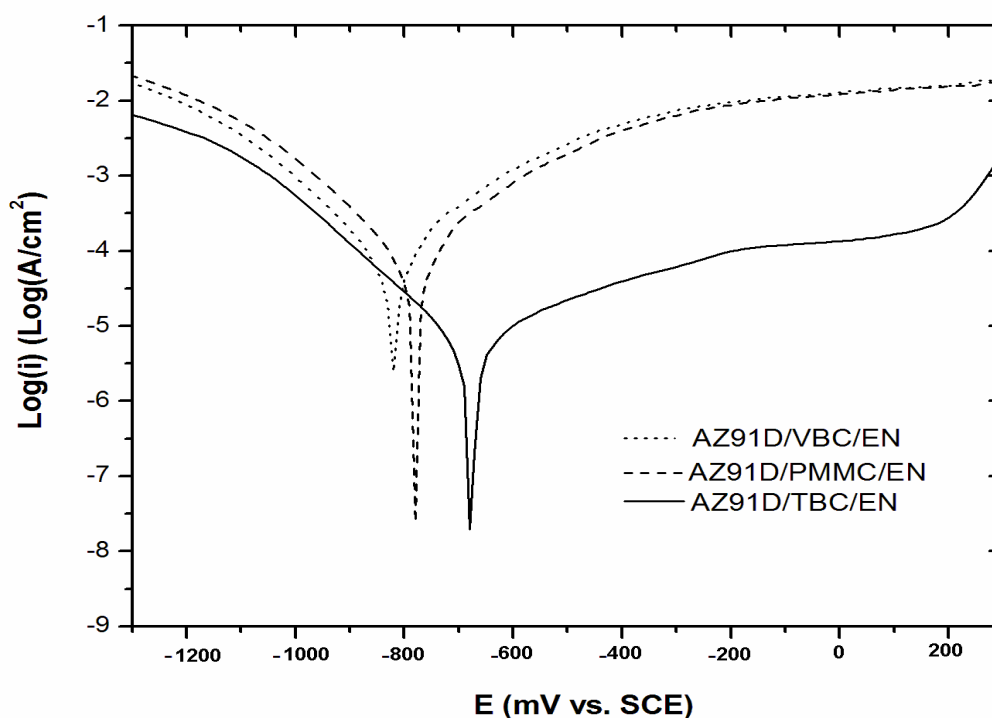


Figure 9. Tafel plots for the EN deposited magnesium alloy with different chromium-free conversion coatings. The experiments were conducted immediately after immersion of the samples in the electrolyte (conducted at room temperature in 3.5 wt% NaCl at a scan rate of 5 mV/s).

Fig. 9 shows Tafel curves recorded in 3.5 wt% NaCl solution for the EN deposited magnesium alloy with different pre-coatings at room temperature. Linear Tafel regions were obtained for Tafel experiments conducted immediately after immersion of the sample in the electrolyte and also freshly prepared samples were used to avoid passivation. However, establishing linear Tafel regions, especially in the anodic portion of the polarization curves (for AZ91D/TBC/EN) was difficult (Fig. 9). Often experimental curves do not display the expected log/linear Tafel behaviour with both anodic and cathodic branches exhibiting curvature [32]. Harvey and Paul developed a non-commercial computer program (SYMADEC) for the interpretation of polarization curves for the Fe/H₂O/H⁺/O₂ corrosion system. The SYMADEC allows for the insertion of different values of film resistance and subsequent calculation of corrosion rate from the polarization curves not exhibiting a Tafel region [32, 33]. The Tafel slopes β_a and β_c were estimated from the polarization curves. Least values of β_a and β_c lead to least corrosion rate according to the Stern-Geary relation [34].

$$i_{corr} = \frac{\beta_a \beta_c}{2.303 \times R_p (\beta_a + \beta_c)} \tag{3}$$

where, *i_{corr}* is the corrosion current density and *R_p* is the polarization resistance. And corrosion rates conversion were calculated by the following equation.

$$1(\text{mA/cm}^2) = \left(\frac{3.28M}{nd} \right) \text{mm/y} \tag{4}$$

where, *M* is atomic mass, *n* is number of electrons freed by the corrosion reaction and *d* is the density. For nickel, 1(mA/cm²) = 10.93 mm/y. However, the nickel is associated with phosphorous content in this electroless NiP deposit, so the value gives approximation only.

Table 7. Results of Tafel extrapolation experiment conducted in 3.5 wt. % NaCl at a scan rate of 5 mV/s. The experiment was conducted immediately after immersion of the samples in the electrolyte.

Sample	β _c	β _a	i _{corr} (Tafel extrapolation)		R _p (Ω cm ²)
			μA/cm ²	mm/y	
AZ91D/VBC/EN	-0.155	0.199	103.87	1.135	2929
AZ91D/PMMC/EN	-0.1467	0.1676	88.07	0.9626	5800

The results of Tafel extrapolation experiment is given in Table-7. Both AZ91D/VBC/EN and AZ91D/PMMC/EN exhibited good Tafel linear region and their corrosion current densities *i_{corr}* were calculated to be 103.87 μA/cm² (1.135 mm/y) and 88.07 μA/cm² (0.9626 mm/y), respectively (table-7).

However, AZ91D/TBC/EN curve not exhibited the expected linear Tafel behaviour in cathodic curvature region to find out the corrosion rate.

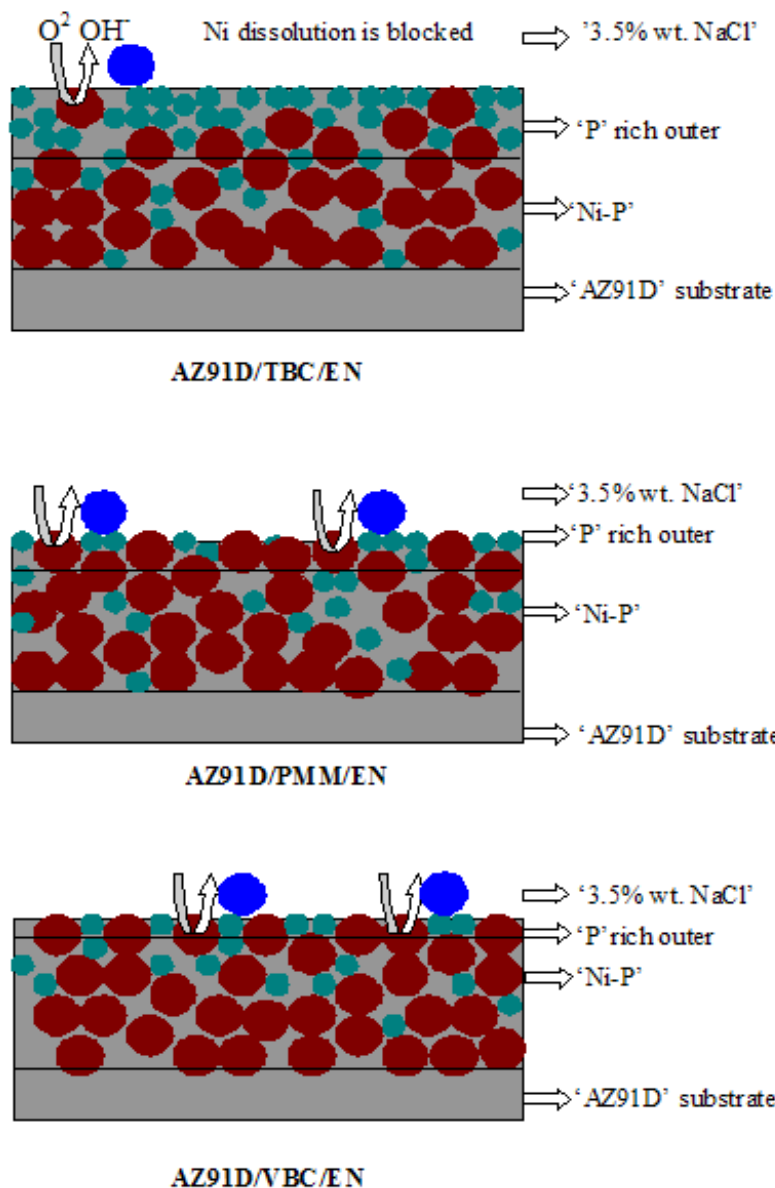
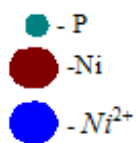
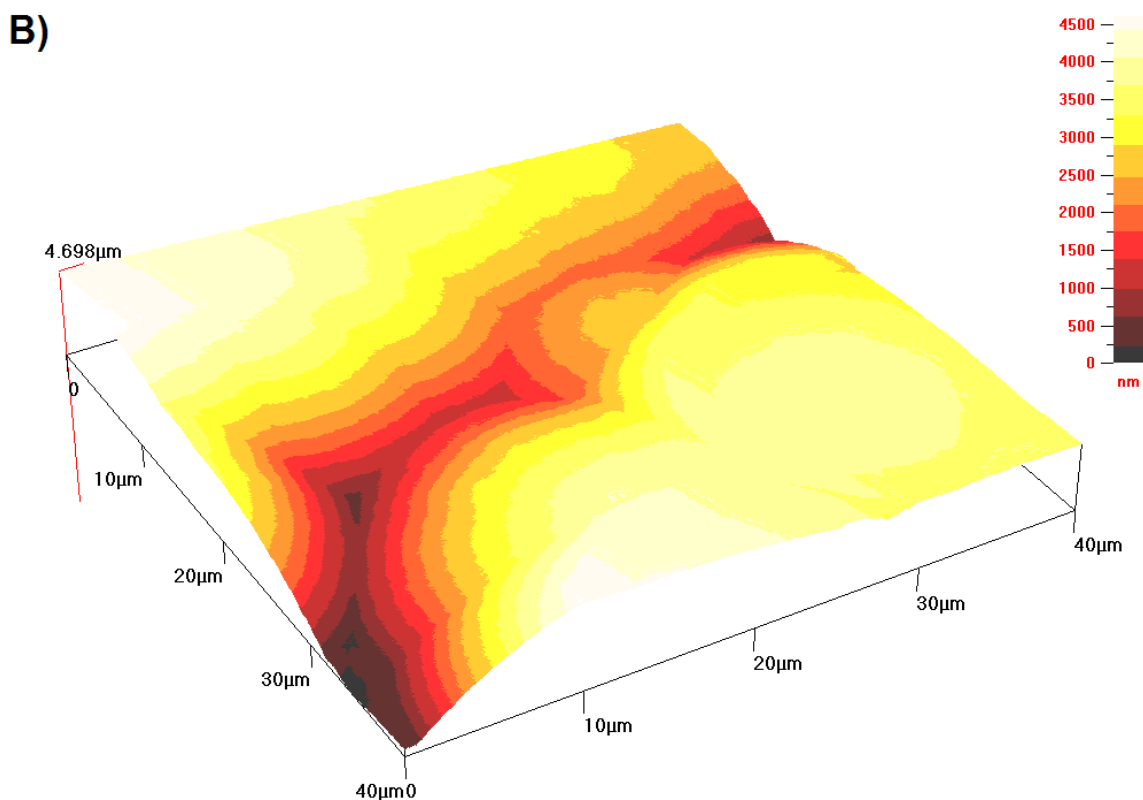
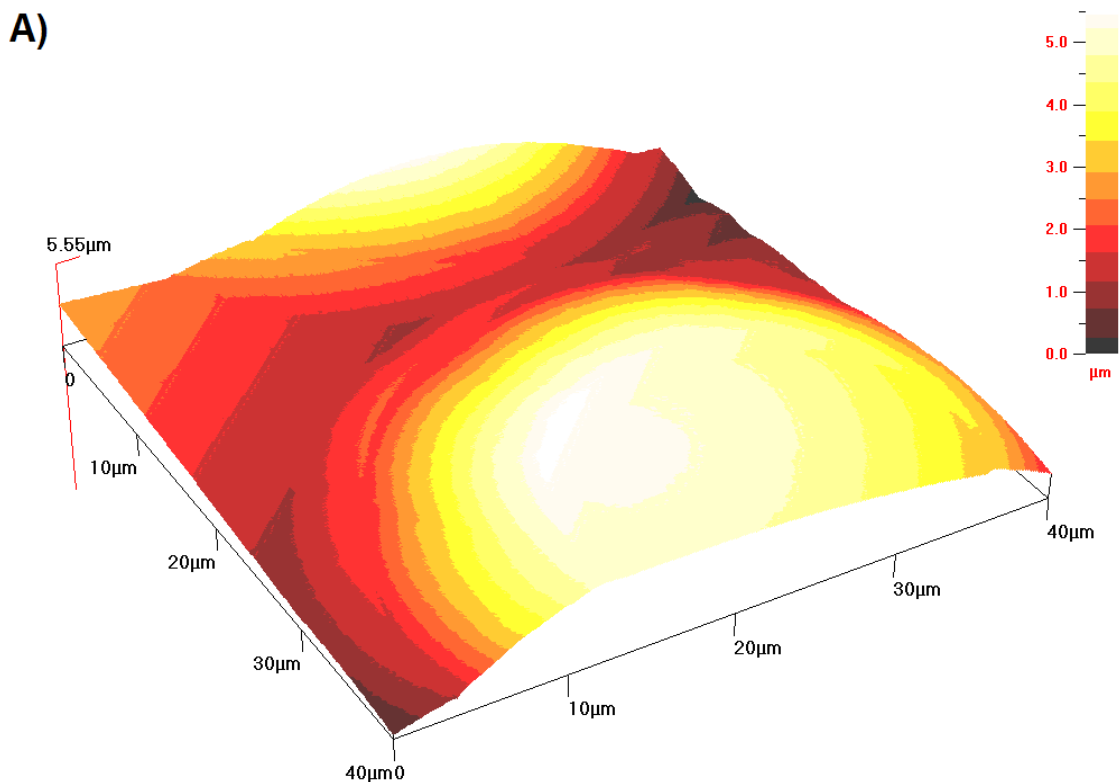


Figure 10. Schematic models for chemical passivation of EN deposits with different chromium-free conversion coatings during corrosion at room temperature in 3.5 wt% NaCl solution.

However, the main aim of this study is to find the performance (corrosion resistance among the different pre-treated EN). From this Tafel plots, one can conclude that the corrosion resistance was

good for TBC compare to other pre-treatments in this study. In addition, the corrosion rate of AZ91D/PMMC/EN was slightly higher compared to the corrosion rates of AZ91D/VBC/EN. Obviously, TBC had relatively higher than these two cases.



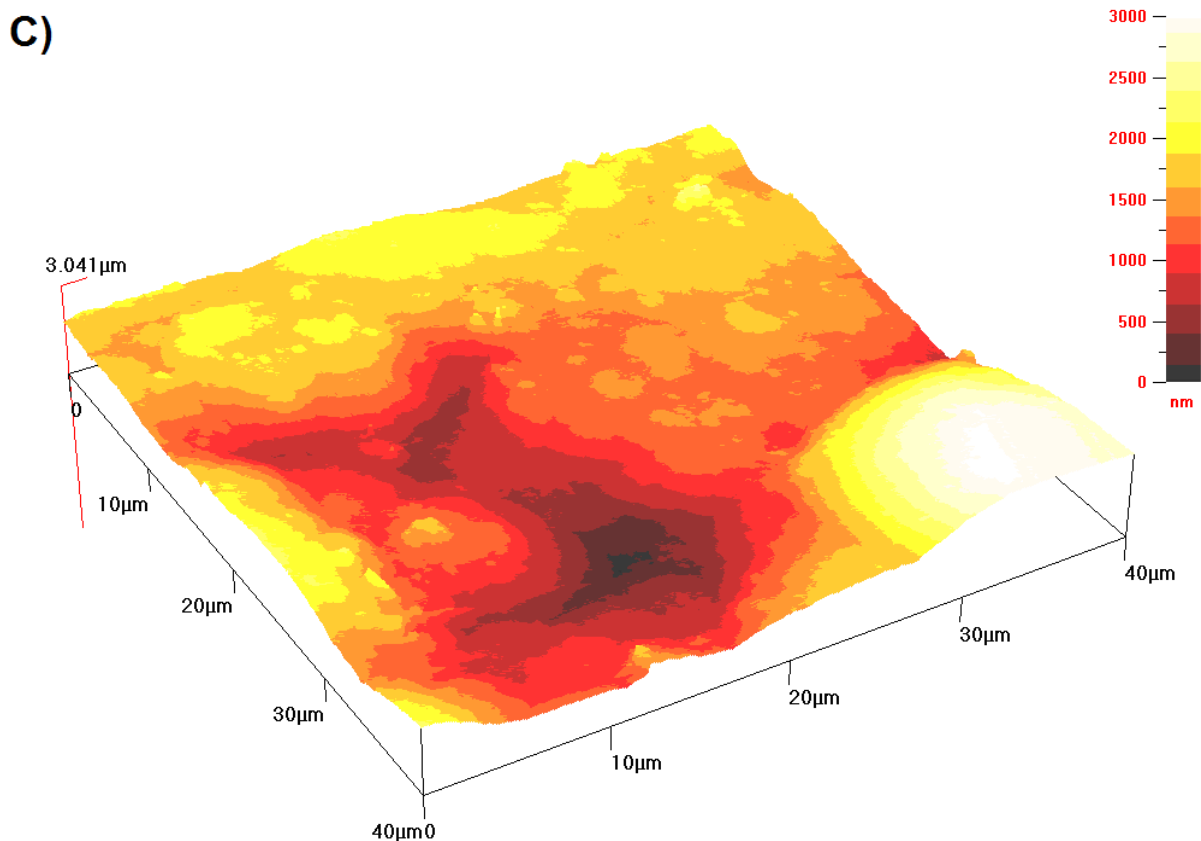


Figure 11. AFM images of EN deposits with: (A) Phosphate-manganese-molybdate; (B) Vanadium based coating; (C) Tannic based coating, on AZ91D alloy.

The schematic diagram (in Fig. 10) illustrated (why TBC/EN had passivation); the general preferential dissolution of nickel produces a P-rich chemically passive layer which hinders further dissolution of nickel in TBC case. The amorphous electroless NiP deposits passivate in acidic and neutral solution, but the presence of crystalline structures increased corrosion rates dramatically [35] and therefore, the present case had good agreement with this. All smooth surfaces possess some degree of roughness, even if only at the atomic level.

Correct function of the fabricated component often is critically dependent on its degree of roughness [36]. The AFM images of electroless NiP deposits with different pre-treatment were presented in Fig. 11 (A, B & C). Fig. 11 (A) & (B) shows surface topography of electroless NiP deposits with VBC and PMMC. The Ni particle deposited on the substrate surface gives rise to peaks and valleys. The average R_a value is about 3.053 and 2.986 μm for VBC and PMMC, respectively. Fig. 11 (C) shows the surface topography of EN deposits with TBC. The Ni deposited on the substrate surface is more and uniform. The AFM image of EN with TBC appears to be relatively smooth and the R_a value is 1.477 μm .

This indicates the surface irregularities were improved to smoothness for TBC than other two cases. Metal cations are more readily dissolved from surface irregularities (surface roughness) than a flat face, since the numbers of bonds on an atom at such an irregularity are fewer and the packing is

more irregular. This also can be evident for the increase in corrosion resistance of TBC pre-treated EN than others and the electrochemical measurement analyses conclude the same.

4. CONCLUSIONS

In this study, the effects of chromium-free pre-treatments on the corrosion properties of electroless NiP deposit were investigated by electrochemical polarization measurements. The main conclusions of this research investigation are:

1. The XPS study reveals the phases of the chromium-free coatings on AZ91D alloy. SEM and EDS reveal that the different chromium-free pre-treatment influencing not only the morphology, but also changed the type ('P' content) of the EN deposits as well. This marginal increase in phosphorus (AZ91D/TBC/EN) forms a layer of adsorbed hypophosphite anions, thereby preventing the hydration of nickel and forming a passive film.

2. All the pre-treated electroless NiP samples exhibited active-passive polarization behaviour. The shift of zero current potential towards positive direction with ascending order (VBC<PMMC<TBC) has been related to modification in cathodic reaction (hydrogen reduction) processes. The passive current densities for TBC/EN were lower than that of other cases and this has been related to the non-defective nature of passive film formed on TBC/EN. The breakdown potential for all electroless NiP was more or less similar in nature. From the Tafel plots, the corrosion resistance was good for TBC/EN compare to other pre-treated electroless NiP deposits.

3. The AFM result shows that the atomic level smoothness was found in the TBC/EN. Metal cations are more readily dissolved from surface irregularities (surface roughness) than a flat face. This was the evident reason for the marginal increase in corrosion resistance of TBC/EN than other pre-treated in this study.

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References

1. B. Landkof, in: K.U. Kainer (Ed.), Proc. Magnesium Alloys and their Applications, Wiley-VCH Verlag, Weinheim, (2000) 168.
2. A.K. Sharma, M.R. Suresh, H. Bhojraj, H. Narayanamurthy, S.P. Sahu, *Met. Finish.* (1998) 10.
3. G.O. Mallory, J.B. Hajdu (Eds.), *Electroless Plating: Fundamentals and Applications*, AESF Publishing, Orlando, FL, (1991) 261.
4. J. Sudagar, J.S. Lian, X.M. Chen, P. Lang, Y.Q. Liang, *Trans. Nonferrous Met. Soc. China.* 21(2011) 921.

5. F.A. Lowenheim, *Modern Electroplating*, Wiley, New York, (1974) 81.
6. H. Umehara, M. Takaya, S. Terauchi, *Surf. Coat. Technol.* 169 (2003) 666.
7. K.Z. Chong, T.S. Shih, *Mater. Chem. Phys.* 80 (2003) 191.
8. M.A. Gonzalez-Nunez, P. Skeldon, G.E. Thompson, H. Karimzadeh, P.Lyon, T.E. Wilks, *Corros. Sci.* 37 (1995) 1763.
9. R.G. Buchheit, S.B. Mamidipally, P. Schmutz, H.Guan, *Corrosion.* 58 (2002) 3.
10. Katya Brunelli, Manuele Dabala, Irene Calliari, Maurizio Magrini, *Corros. Sci.* 47 (2005) 989.
11. L. Rudd, C.B. Breslin, F. Mansfeld, *Corros. Sci.* 42 (2000) 275.
12. Xiaoming Chen, Guangyu Li, Jianshe Lian, Qing Jiang, *Surf. Coat. Technol.* 204 (2009) 736.
13. H.J. Kim, J. Zhang, R.H. Yoon, R. Gandour, *Surf. Coat. Tech.* 188 (2004) 762.
14. J.S. Lian, G.Y. Li, L.Y. Niu, C.D. Gu, Z.H. Jiang, Q. Jiang, *Surf. Coat. Technol.* 200 (2006) 5956.
15. G.Y. Li, J.S. Lian, L.Y. Niu, Z.H. Jiang, *ISIJ Int.* 45 (9) (2005) 1326.
16. K.H. Yang, M.D. Ger, W.H. Hwu, Y. Sung, Y.C. Liu, *Mater. Chem. Phys.* 101 (2007) 480.
17. J. Sudagar, Y.Q. Liang, G.Y. Li, Q. Jiang, J. S. Lian, *Mater. Chem. Phys.* (submitted).
18. G. L. Song, A. Atrens, M. Dargusch, *Corros. Sci.* 41 (1998) 249.
19. G. L. Song and A. Atrens, *Adv. Eng Mater.* 1 (1999) 11.
20. G. Mattoigno, C. Ferragina, M.A. Massucci, P. Patrono, A. La Ginestra, *J. Electron Spectrosc. Relat. Phenom.* 46 (1988) 285.
21. D.T. Clark, W.J. Feast, P.J. Tweedale, H.R. Thomas, *J. Polym. Sci: Polym. Chem. Ed.* 18 (1980) 1651.
22. T. Yoshida, K. Yamasaki, S. Sawada, *Bull. Chem. Soc. Jpn.* 51 (1978) 1561.
23. C.D. Wagner, *J. Electron Spectrosc. Relat. Phenom.* 18 (1980) 345.
24. R. Hoogewijs, L. Fiermans, J. Vennik, *J. Electron Spectrosc. Relat. Phenom.* 11 (1977) 171.
25. C. Malitesta, G. Razzini, L. Peraldo bicelli, L. Sabbatini, *Int. J. Hydrogen Energy.* 12 (1987) 219.
26. D. Borgmann, E. Hums, G. Hopfengartner, G. Wedler, G.W. Spitznagel, I. Rademacher, *J. Electron Spectrosc. Relat. Phenom.* 63 (1993) 91.
27. Liping Wang, Junyan Zhang, Yan Gao, Qunji Xue, Litian Hu, Tao Xu, *Scripta Mater.* 55 (2006) 657.
28. P.H. Lo, W.T. Tsai, J.T. Lee, M.P. Hung, *Surf. Coat. Technol.* 67 (1994) 17.
29. T.S.N. Sankara Narayanana, I. Baskaranb, K. Krishnavenia, S. Parthiban, *Surf. Coat. Technol.* 200 (2006) 3438.
30. Z. Guo, K.G. Keong, W. Sha, *J. Alloys Compd.* 358 (2003) 112.
31. Woo-Jae Cheong, Ben L. Luan, David W. Shoesmith, *Corros. Sci.* 49 (2007) 1777.
32. Harvey J. Flitt, D. Paul Schweinsberg, *Corros. Sci.* 47 (2005) 3034.
33. H.J. Flitt, D.P. Schweinsberg, *Corros. Sci.* 47 (2005) 2125.
34. M. Stern, A.L. Geary, *J. Electrochem. Soc.* 104 (1957) 56.
35. A.Krolkowski, B. Pokrywa, *Electrochem. Acta.* 38 (1993) 1979.
36. R. Elansezhian, B. Ramamoorthy, P. Kesavan Nair, *J. Mater. Process Tech.* 209 (2009) 233.