

**Effect of Zn content on corrosion behavior of Mg-Y-Zn alloys**

Y. Shi,° X.-Q. Liu,°,*, Z.-L. Liu,°, H.-J. Xie,° Y.-H. Wang,°, J. Li,°,b,c

°College of Materials Science and Technology, Nanjing University of Aeronautics and Astronautics, Nanjing, Jiangsu, China

b Jiangsu (Zhongyi) Engineering Research Center of Light Alloy Precision Die Casting, Changshu, Jiangsu, China
c Jiangsu Favour Automotive New Stuff Sci-Tech Co., Ltd., Changshu, China

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Abstract

The microstructure, corrosion behavior and electrochemical behavior of as-cast Mg-4Y-xZn (x=1,2,3,4 wt.%) are studied by SEM, weight loss and electrochemical tests. Mg_{12}Y_{2}Zn (X), Mg_{3}Y_{2}Zn_{3} (W) and Mg_{24}Y_{5} constitute the phase composition system of the alloy. When Zn content is 1 wt.%, all tests reveal that alloy has the optimal corrosion performance. The second phase in these alloys, due to their nobler nature than α-Mg, exists as cathode during corrosion process, so that α-Mg preferentially occurs corrosion to accelerate the formation of corrosion pits. After soaking in 3.5 wt.% NaCl solution for some time, the stability of the W phase changed, and gradually dissolved, which was finally removed by chromic acid used for removal of corrosion products. In addition, X phase can be used as an anode in the micro-galvanic cells formed with W phase to reduce the corrosion rate of α-Mg and thus improve the corrosion performance of the alloy.

Keywords: Mg-Y-Zn; second phase; corrosion behaviors; electrochemical behaviors

1. Introduction

As the most promising lightweight and green engineering material, magnesium alloy has the advantages of light specific gravity, high specific strength, good damping performance and easy recycling, making it has a wide range of application prospects in the electronic communications, aerospace and other fields [1-5]. However, the development of magnesium alloy is hindered due to its low absolute strength and poor corrosion resistance [6, 7].

Recently, Mg-Y-Zn series alloys have aroused a lot of attention because of their excellent performance [8-13]. Furthermore, the effect of LPSO structure on relative corrosion performance and how to enhance the corrosion resistance are currently being explored. Zhang et al. revealed corrosion pits were mostly concentrated at the junction of LPSO phase and α-Mg, which reflected that LPSO phase accelerated the micro-galvanic corrosion of α-Mg [14]. C. Q. Li et al. found that LPSO with different volume fraction affected the corrosion performance of the alloy. The dense and thick LPSO phase hindered the corrosion expansion to a certain extent. The distribution of LPSO phase changed the morphology of corrosion pits by affecting the shape of magnesium dendrite [15]. Lei Bao et al. also proposed that the species of the phase in the alloy affected the corrosion resistance and mechanism. W phase causes severe intergranular corrosion.

Corresponding Author:
E-mail address: liuxiqin@nuaa.edu.cn
corrosion and forms micropores. Dense lath LPSO phase can effectively protect the matrix \cite{16}. Similar results have been confirmed in many subsequent studies \cite{17-22}.

Based on the above research, it mainly covers the influence of LPSO phase content and morphology on the corrosion performance and process of magnesium alloy, while W phase is not conducive to the mechanical and corrosion properties of magnesium alloy. However, recent studies have shown that when LPSO and W phase coexist in the alloy, they have synergistic effect on the mechanical performance of the alloy \cite{23}. At present, there are few researches on the corrosion properties of the magnesium alloy with LPSO and W phase, so it is necessary to study its mechanism.

Therefore, based on the current research background, the relationship between the corrosion behavior of the phase and the $\alpha$-Mg needs to be further investigated. In this paper, Mg-4Y-xZn ($x=1,2,3$ and 4 wt.%) alloys with different second phases and their contents were studied with the aim of elucidating the relationship between the phases and the corrosion mechanism of the alloys.

2. Materials and methods

2.1. Alloys preparation and microstructural analysis

The studied alloys were manufactured by melting pure Mg, pure Zn and Mg-30\% Y master alloys. Table 1 shows the chemical compositions of the studied alloys. The protective gas is a mixture atmosphere in which the volume ratio of SF$_6$ to CO$_2$ is 1:100. It was smelted in a steel crucible at about 720 $^\circ$C, stirred at 740 $^\circ$C for 30 min, and then cast in a preheated (~ 200 $^\circ$C) iron mold at about 720 $^\circ$C. The ingot size was 25×100×150 mm.

| Alloys | Y (wt.\%) | Zn (wt.\%) | Mg (wt.\%) |
|--------|-----------|------------|------------|
| WZ41   | 4         | 1          | Balance    |
| WZ42   | 4         | 2          | Balance    |
| WZ43   | 4         | 3          | Balance    |
| WZ44   | 4         | 4          | Balance    |

The samples were mechanically grinded by SiC paper (from 120 grit to 3000 grit), then mechanically polished with 0.5 $\mu$m diamond paste. The X-ray diffractometer (XRD, MXP2NAHF) taking scanning range of 10$^\circ$-90$^\circ$ with increments of 20 and scanning rate of 5$^\circ$/min with Cu K$\alpha$ radiation at 40 kV and 150 mV which was used for phase identification of the samples and corrosion products. The microstructure, intermetallic compounds and corrosion morphology of the specimens were observed by scanning electron microscopy (SEM, LYRA3TESCAN). The phase compositions were analyzed by the energy dispersive spectroscopy (EDS) equipped in SEM.

2.2. Immersion tests and mass loss tests

For the purpose of explore the corrosion properties of different samples, the soaking experiment was taken in 3.5 wt.% NaCl solution at 25 $^\circ$C for a certain time. Before immersion, the specimens were grinded by SiC papers down to 2000 grit. The ratio of the surface area of the specimen to the volume of NaCl solution was controlled at 1:100 (exposed surface to the solution was 6 cm$^2$ to 600 ml). After immersion, the sample was
washed in a hot mixture of 200 g/L CrO$_3$+10 g/L AgNO$_3$ to remove the corrosion products on the alloy, and then cleaned with distilled water and desiccated in the oven. Then, the corrosion morphology of the corroded specimens was observed by SEM. The actual corrosion rate of the alloy is evaluated by the mass loss before and after the removal of corrosion products, and is calculated by the following formula:

$$CR = \frac{8.76 \times 10^4 \times \Delta g}{A \times t \times \rho}$$  \hspace{1cm} (1)

Where $\Delta g$ (g) is the mass loss, $A$ (cm$^2$) is the surface area of the specimen, $t$ (h) is the soak time and $\rho$ (g/cm$^3$) is the alloy density. Three experiments were carried out on the four alloys and the average value were applied.

2.3. Electrochemical tests

In 3.5 wt.% NaCl solution, the alloy was electrochemically tested by standard three electrode cell in Zahner Zennium electrochemical workstation. Among them, four specimens acted as working electrode, a platinum electrode acted as the counter electrode, a saturated calomel electrode (SCE) acted as reference electrode. All specimens were sealed with epoxy resin plastic, 1 cm$^2$ exposed surface was used as working electrode surface, and the other end was connected with copper wire. EIS measurement was taken in the frequency range of 100 kHZ-10 mHZ, using 10 mv sinusoidal voltage, the experimental results were fitted by ZsimpWin. Potentiodynamic scanning was carried out at the rate of 1 mv/s, the limit of polarization current was -100 mA for cathode and 100 mA for anode. The above experiments were carried out after the samples have stabilized at open circuit potential (OCP) for 30 min.

3. Results and Discussion

3.1. Phase constitutions and microstructure observation

Fig. 1. XRD spectrums of as-cast Mg-4Y-xZn alloys

Fig. 1 shows the XRD spectrums of the Mg-4Y-xZn alloys, indicating that when the Zn content exceeds 2 wt. %, the content of Mg$_{24}$Y$_5$ phase decreased until disappeared,
while the W phase began to appear and the content gradually increased. In addition to the α-Mg, WZ41 is composed of X phase and Mg$_{24}$Y$_5$, WZ42 and WZ43 are both composed of X phase, W phase and Mg$_{24}$Y$_5$, and WZ44 is composed of W phase$^{[24, 25]}$.

Fig. 2 shows the SEM microstructure and EDS results of Mg-4Y-xZn alloys. WZ41 is composed of matrix, lamellar phase and blocky white granules. WZ42 and WZ43 are composed of matrix, lamellar phase, blocky white granules and WZ44 is only composed of matrix and white eutectic network. With the increase of zinc content, the lamellar phase gradually decreases, while the reticular white eutectic phase gradually increases. Combining the results of XRD, EDS and the detailed analysis in our previous work$^{[12]}$, it can be obtained that the white massive particles corresponding to point A are Mg$_{24}$Y$_5$, the flake phase corresponding to point B is X phase, the white eutectic network corresponding to point C is W phase, and the matrix is α-Mg. According to Fig. 2(d), W phase in WZ42 appears in the lamellar X phase as a sheet. Fig. 2(f) shows that the W phase in WZ43 appears near the surface of X phase$^{[26-28]}$.  

**Fig. 2.** SEM morphologies of as-cast Mg-4Y-xZn alloys: (a) and (b) WZ41, (c) and (d) WZ42, (e) and (f) WZ43, (g) and (h) WZ44, (i) EDS result of point A in (b), (j) EDS result of point B in (d), (k) EDS result of point C in (h)
3.2. Corrosion behaviors

Fig. 3. Average corrosion rate calculated from the weight loss after immersion in 3.5 wt. % NaCl solution at 25 °C for 72 h

Fig. 3 is the average corrosion rate of as-cast Mg-4Y-xZn alloys immersed in 3.5 wt.% NaCl solution at 25 °C for 72 h. The results reveal that the corrosion rate increases with the increase of Zn content. Moreover, the corrosion rate of WZ41 is the lowest, and that of WZ44 is the highest. Nevertheless, WZ43 shows lower corrosion rate than WZ42, combine with the microstructure of the alloy, this may be because WZ43 alloy contains more W phase than WZ42 alloy, which affects the corrosion rate.

Fig. 4. Corrosion surface after immersion in 3.5 wt.% NaCl solution for 12 h with removal of corrosion products:
(a) WZ41; (b) WZ42; (c) WZ43; (d) WZ44

The corrosion surface of the alloy immersed for 12 h and removed the corrosion products is shown in Fig. 4. Obviously, the corrosion began and gradually expanded in
the form of pitting corrosion, and a certain number of corrosion pits appeared on the alloy surface. Among them, WZ41, WZ42 and WZ43 alloys showed the characteristics of filamentous corrosion, part of uncorroded substrate still exists in the surface of the above alloys, demonstrating the relatively low corrosion pit expansion rate and weak corrosion occurs. With the increase of Zn content, the characteristics of filamentous corrosion became insignificant [29]. The surface of WZ44 alloy shows the most serious corrosion behavior. The bare matrix is almost invisible. The surface is composed of corrosion pits with different shapes and sizes.

Fig. 5. Corrosion micrographs of (a-d) WZ41 and (e-h) WZ42 after immersion in 3.5 wt.% NaCl solution for (a, e) 5 min, (b, f) 30 min, (c, g) 2 h, (d, h) 12 h and removal of the corrosion products

Fig. 6. Corrosion micrographs of (a-d) WZ43 and (e-h) WZ44 after immersion in 3.5 wt.% NaCl solution for (a, e) 5 min, (b, f) 30 min, (c, g) 2 h, (d, h) 12 h and removal of the corrosion products

SEM images shown in Fig. 5 and 6 reveal the corrosion morphology of the alloy after immersion for 5 min, 30 min, 2 h and 12 h. For WZ41 alloy, the corrosion pits first appear in the vicinity of X phase and Mg$_2$Y$_5$ phase, while the center of $\alpha$-Mg and second phase is basically intact, indicating that the corrosion begins at the junction of $\alpha$-Mg and second phase. Then the corrosion pits gradually extended to the $\alpha$-Mg, and part of the second phase gradually detach from the surface due to the loss of support from matrix. And then, the second phase is almost completely detached; Finally, it shows that the corrosion pits have begun to expand to the matrix and show the characteristics of filamentous corrosion. For WZ42 and WZ43 alloys, similar to WZ41
alloy, the corrosion pits begin at the junction of the second phase and matrix. Then the X phase was dissolved or detached, but the W phase was retained; And then, the second phase was completely removed, the corrosion pits began to expand to matrix and left obvious corrosion pits; Finally, the corrosion pits continue to expand and have bare matrix residues, showing the characteristics of filamentous corrosion. For WZ44 alloy, the corrosion pits begin at the junction of W phase and matrix; Then the W phase was gradually eroded and removed leaving a reticular gully; After that, the corrosion pits began to expand to the matrix, leaving only some uncorroded matrix; Finally, the alloy surface is completely occupied by corrosion pits, and there is no exposed matrix. In summary, it can be found that the corrosion pits of the four alloys began at the junction of the second phase and the $\alpha$-Mg, and then the second phase was gradually eroded or detached, and then began to expand to the matrix, and finally formed obvious corrosion pits. However, for WZ42 and WZ43 alloys, after immersion for 30 min, it can be found that W phase is retained compared with X phase. Considering that the W phase still exists, the possibility of the X phase falling off is not very great, which may be caused by the formation of micro-galvanic cells between X phase and W phase leading to the dissolution of the X phase as an anode. As shown in Fig. 7, the EDS results for point D in Fig. 6 (b) also confirm that the W phase is retained during the initial corrosion process, but a slight corrosion occurs on its surface and is gradually eroded during the subsequent corrosion expansion. As for four alloys, in addition to WZ44 alloy, WZ41, WZ42 and WZ43 alloy eventually showed the characteristics of filamentous corrosion; Secondly, the corrosion pit expansion rate of WZ41 alloy is the lowest, and that of WZ44 alloy is the highest, which has been seriously expanded to matrix at 2 h.

Previous studies have shown that the second phase in Mg alloy can present as cathode in the corrosion process to accelerate micro-galvanic corrosion. Siqi Yin et al. revealed that compared with the matrix, W phase and X phase have higher potential, and W phase is electrochemically nobler than X phase [30, 31]. Therefore, the type, quantity of the second phase have a significant impact on the corrosion process [32, 33]. Obviously, the micro-galvanic corrosion occurs at the junction of the second phase with $\alpha$-Mg, which makes the $\alpha$-Mg dissolve as an anode. With the extension of soak time, the corrosion pits gradually expanded, and then the second phase eroded or detached to form corrosion ravine. A. Srinivasan et al. also confirmed that sodium chloride solution
changed the stability of the W phase at the end of corrosion, and the corrosion products strongly alkalized the surface after long-term immersion, so that the exposed W phase gradually dissolved and removed \textsuperscript{34}. Similar observations were made by Neil et al. in ZE41 alloy \textsuperscript{35,36}. In addition, Ziyou Ding et al found that during the corrosion process of Mg-Li-Ca alloy, the second phase will occur exfoliation corrosion under the synergistic effect of pitting, filamentous and intergranular corrosion, and the second phase will fall off from the matrix \textsuperscript{37}. Guijia Gao et al also obtained similar conclusions \textsuperscript{38}. As for WZ41 alloys, due to the low content of X phase, the number of micro-galvanic cells formed is limited, which shows the best corrosion performance. As for WZ42 and WZ43 alloys, due to the X phase is stable in the sodium chloride solution and will not be dissolved, but W phase is electrochemically nobler than X phase. Therefore, micro-galvanic cells will may be formed at the junction of X phase and W phase, so that the X phase is used as an anode to dissolve during the corrosion process, thereby reducing the number of micro-galvanic cells formed between the second phase and matrix, whereupon slowing down the corrosion rate of matrix to some extent. Since WZ43 alloy contains more W phase which is located on the surface of X phase, more X-W micro-galvanic cells are formed, so it has better corrosion resistance than WZ42 \textsuperscript{38,39}. The results of the above-mentioned weight loss experiment also verify this result. In addition, in the later stage of corrosion development, some X phase will also fall off from the matrix due to exfoliation corrosion, which is also observed in Fig. 5 and 6. As for WZ44 alloy, due to the higher potential of W phase than α-Mg, the trend of micro-current corrosion is the largest, showing more obvious corrosion characteristics and the worst corrosion performance. For Mg\textsubscript{2}Y\textsubscript{x} phase, although micro-galvanic cells can also be formed with α-Mg, it has little effect on the overall corrosion behavior due to its low content and detached at the end of corrosion.

3.3. Electrochemical testing

Fig. 8. Potentiodynamic polarization curves of WZ41, WZ42, WZ43 and WZ44 alloys measured in 3.5 wt.% NaCl solution

Fig. 8 shows the polarization curves of Mg-4Y-xZn alloy in 3.5 wt.% NaCl solution, the results fitted by Tafel extrapolation method and listed in Tabe 1. It is generally
believed that the cathode branch is related to the reactive precipitation of hydrogen, while the anode branch is related to the reactive dissolution of matrix. It can be seen that the cathode branch and anode branch in the polarization curve are obviously different, and the increase rate of the curve in the anode branch is much larger than that in the cathode branch [40-42]. In addition, from the cathode point of view, the alloy system has similar polarization behavior and a wide linear Tafel region, which indicates that the alloy has uniform and stable corrosion rate. As for the anode branch, no obvious passivation phenomenon was found, reveal that the phase composition of the alloy had little effect on the formation and dissolution of the surface oxide film [34]. According to the fitting results of table 2, the corrosion current density ($J_{\text{corr}}$) of the four alloys is WZ44 > WZ42 > WZ43 > WZ41, and the polarization resistance ($R_p$) increases in turn. It is generally believed that the value of $J_{\text{corr}}$ reflects the actual corrosion rate of the alloy from corrosion kinetics, so WZ41 alloy shows the best corrosion performance. However, WZ43 alloy has the most negative corrosion potential ($E_{\text{corr}}$). From the thermodynamic point of view, the corrosion tendency of WZ43 alloy is the highest, but actually it does not show the worst corrosion performance. This also reflects that the actual corrosion rate and mechanism of the alloy are closely related to the microstructure and other factors [30].

| Alloys | $E_{\text{corr}}$ /V | $J_{\text{corr}}$ /μA·cm$^{-2}$ | $\beta_a$ /mv·dec$^{-1}$ | $\beta_c$ /mv·dec$^{-1}$ | $R_p$ /Ω |
|--------|-----------------|-----------------|-----------------|-----------------|--------|
| WZ41   | -1.52           | 130             | 45.4            | -292            | 131    |
| WZ42   | -1.53           | 231             | 48.1            | -298            | 77.9   |
| WZ43   | -1.55           | 221             | 58.8            | -308            | 96.8   |
| WZ44   | -1.53           | 289             | 55.7            | -529            | 75.8   |

Fig. 9. Electrochemical impedance spectrum (EIS) plots of the alloys: (a) Nyquist plots; (b) Bode phase angle and impedance plots

Fig. 9 shows the EIS curves measured after the 3.5 wt.% NaCl solution was stabilized for 3600 s. In Nyquist plot, the Mg-4Y-xZn alloys show obvious characteristics of high frequency capacitance ring and low frequency inductance ring. Generally, the former can be attributed to the charge transfer process of the double layer and the electrode/electrolyte interface, while the latter can be ascribed to the damage of the surface film and the corrosion nucleation at the incipient stage of local corrosion. Among them, the larger the radius of the high frequency capacitance ring, the larger the
charge transfer resistance, which reflects the stability of the surface layer, so that the impedance is significantly improved \[^{[41, 42]}\]. With the increase of Zn content, the size of the capacitance ring is roughly reduced, indicating that the corrosion resistance is gradually reduced. However, WZ43 alloy shows a larger capacitance ring than WZ42 alloy, which is also caused by the formation of X-W micro-galvanic mentioned above. In addition, all alloys show certain induction characteristics, among which WZ42 is the most obvious, which means that the surface film of WZ4 alloy is very unstable and has poor protection. The Bode phase angle and impedance plots are shown in Fig. 9(b), as for the Bode plots of phase angle(θ) vs. log \( f \), it can be seen that the four alloys are composed of a peak and a trough, usually, a peak represents a time constant. In this case, the peak represents the high-frequency capacitance ring, and the trough represents the low-frequency inductance ring. The large phase angle peak and peak width in the WZ41 alloy demonstrate large resistance and small capacitance, so as to the preferable corrosion protection. As for the Bode plots of log \( |Z| \) vs. log \( f \), in the high frequency region, the impedance values of the four alloys increased significantly, the increase rate slowed down and gradually stabilized in the intermediate frequency range, and the impedance value began to decrease in the low frequency region. Among them, with the increase of Zn content, the variation trend of the impedance modulus of the alloy is similar to that of the capacitance ring size, showing a gradual decrease. The impedance modulus of WZ41 alloy is the largest, which means that it has high polarization resistance. The impedance modulus of WZ43 alloy is larger than that of WZ42, and the specific reason is the same as that of Nyquist. In addition, it can be seen that the impedance modulus of all alloys decreased significantly in the low frequency region, and the impedance modulus of WZ42 alloy decreased fastest, indicating the instability of its surface film.

Fig. 10. Equivalent circuit of the EIS spectra for the alloys immersed in 3.5 wt.% NaCl solution

For reveal the corrosion mechanism of the alloy, the EIS equivalent circuit model shown in Fig. 10 is provided to fit the corrosion system. \( R_s \) is the solution resistance; \( R_{ct} \) is charge transfer resistance; \( R_f \) is membrane resistance. CPE is a constant phase element. Due to the instability of the electrochemical system, the ideal capacitance is replaced by the ideal capacitance to represent the incompleteness of the double layer capacitance. It is defined by two parameters \( Y \) and \( n \). \( Y \) represents the non-ideal capacitance, which is attributed to the dispersion effect caused by cracks, surface oxide films, impurities and the second phase. \( n \) is the dispersion coefficient. When \( n = 1 \), CPE is the same as the pure capacitance, and when \( n = 0 \), CPE is the same as the pure resistance. In addition, \( R_{ct} \) and CPE\(_{dl} \) are combined to describe the high-frequency
capacitor loop; RL represents the resistance, and L represents the inductance, which
together represent the low-frequency inductance loop, showing the beginning of local
corrosion; \( R_f \) and CPE\(_f\) are combined to represent the mass transfer during the
dissolution of corrosion products \([30,31]\).

Table 3 The fitted EIS data on the basis of the equivalent circuits presented in Fig. 10

| Alloys | \( R_S \)  | \( R_{ct} \)  | CPE\(_{dl} \) | \( R_L \)  | L  | \( R_f \)  | CPE\(_f\) |
|--------|------------|------------|-------------|------------|----|------------|-------------|
|         | \( \Omega \cdot \text{cm}^2 \) | \( \Omega \cdot \text{cm}^2 \) | \( Y_{dl}/\Omega \cdot \text{cm}^2 \cdot \text{s}^n \) | \( \Omega \cdot \text{cm}^2 \) | \( \Omega \cdot \text{cm}^2 \) | \( \Omega \cdot \text{cm}^2 \) | \( Y_f/\Omega \cdot \text{cm}^2 \cdot \text{s}^n \) |
| WZ41   | 8.602      | 9.029      | 3.172×10^{-5} | 0.867      | 1945 | 20980      | 331.1       | 1.119×10^{-5} | 1          |
| WZ42   | 9.209      | 6.967      | 1.572×10^{-5} | 0.933      | 308.7 | 1969       | 230.9       | 2.898×10^{-5} | 0.929      |
| WZ43   | 9.005      | 13.12      | 3.051×10^{-5} | 0.882      | 1231 | 12490      | 285.8       | 1.588×10^{-5} | 0.96       |
| WZ44   | 8.411      | 6.785      | 2.219×10^{-5} | 0.925      | 1282 | 8567       | 226         | 2.656×10^{-5} | 0.933      |

The EIS curves measured in the experiment were fitted based on the proposed
equivalent circuit model, and the fitting results are listed in Table 3. Generally speaking,
the higher the values of \( R_{ct} \) and \( R_f \), the lower the dissolution rate of the matrix, the better
the protective performance of the surface film and the better the corrosion resistance.
In general, the \( R_{ct} \) and \( R_f \) values of the Mg-4Y-xZn alloys are basically WZ41 > WZ43 >
WZ42 > WZ44. However, combined with the previous experimental results, WZ41
alloy should have the maximum \( R_{ct} \) value, but it is actually less than WZ43 alloy. This
may be stem from the formation of micro-galvanic cells between W phase and X phase
in WZ43 alloy, which reduces the corrosion rate of matrix at the initial stage of
corrosion \([40]\).

Fig. 11. XRD patterns of corrosion product of Mg-4Y-xZn alloys after immersion in 3.5 wt.% NaCl solution for 72
h at 25 °C.

Fig. 11 shows the XRD spectrums of the corrosion products of the Mg-4Y-xZn
alloys soaked in 3.5 wt.% NaCl solution at 25 °C for 72 h, which reveals that the
corrosion products are consisted of Mg(OH)\(_2\), with the exception of the WZ44 alloy,
the other three alloys also have a small amount of X phase. When the alloy is placed in
the NaCl solution, the reaction is shown as follows:

\[
\text{Mg} + 2\text{H}_2\text{O} \rightarrow \text{Mg(OH)}_2 + \text{H}_2 \tag{2}
\]
According to the previous results, since the content of W phase gradually increases, during the corrosion stage, the X phase is dissolved as an anode due to the formation of X-W micro-galvanic, so that the peak strength of X phase in the corrosion product of WZ41 alloy is the highest. The Mg(OH)$_2$ was formed in the whole corrosion process, and a small amount of X phase can be attributed to the exfoliation corrosion occurring at the grain boundaries of X phase and matrix, subsequently, some small X phases are retained in the corrosion products due to the absence of matrix support. However, the W phase dissolved gradually in the corrosion process, and no W phase was found in the corrosion products.

4. Conclusions

By studying the microstructure, corrosion behavior and properties of Mg-4Y-xZn alloy with different second phase composition in 3.5 wt.% NaCl solution for different soaking time, the following conclusions can be drawn:

(1) With the increase of Zn content, the main second phase in the Mg-4Y-xZn alloy gradually changed from X phase to X + W phase and finally all became W phase;

(2) The immersion test and electrochemical test confirmed that with the increase of Zn content, the corrosion resistance of Mg-4Y-xZn alloy generally shows a gradual decline, but WZ43 alloy showed better corrosion resistance than WZ42;

(3) The type and content of the second phase play an essential role in the corrosion mechanism and corrosion rate of the alloy. The potential between the $\alpha$-Mg and the second phase is different. In the immersion process, the second phase acts as the cathode, and the micro-galvanic corrosion dominates, accelerating the dissolution of the matrix. X phase can be used as an anode in the micro-galvanic cells formed with W phase, which reduced the number of micro-galvanic cells between X phase and $\alpha$-Mg, thereby protecting the matrix from corrosion;

(4) The corrosion products of the Mg-Y-Zn alloy consist mainly of Mg(OH)$_2$. Except for the WZ44 alloy, a small amount of X phase is also present in the corrosion products of the remaining alloys.

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Fig. 1. XRD spectrums of as-cast Mg-4Y-xZn alloys

Fig. 2. SEM morphologies of as-cast Mg-4Y-xZn alloys: (a) and (b) WZ41, (c) and (d) WZ42, (e) and (f) WZ43, (g) and (h) WZ44

Fig. 3. Average corrosion rate calculated from the weight loss after immersion in 3.5 wt. % NaCl solution at 25 °C for 72 h

Fig. 4. Corrosion surface after immersion in 3.5 wt.% NaCl solution for 12 h with removal of corrosion products: (a) WZ41; (b) WZ42; (c) WZ43; (d) WZ44

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