Acceleration of Plasma Nitriding at 550 °C with Rare Earth on the Surface of 38CrMoAl Steel

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Abstract: In order to explore the effect of the addition of rare earth (RE) to a steel microstructure and the consequent performance of a nitrided layer, plasma nitriding was carried out on 38CrMoAl steel in an atmosphere of NH3 at 550 °C for 4, 8, and 12 h. The modified layers were characterized using an optical microscope (OM), a microhardness tester, X-ray diffraction (XRD), a scanning electron microscope (SEM), a transmission electron microscope (TEM), and an electrochemical workstation. After 12 h of nitriding without RE, the modified layer thickness was 355.90 µm, the weight gain was 3.75 mg/cm², and the surface hardness was 882.5 HV0.05. After 12 h of RE nitriding, the thickness of the modified layer was 390.8 µm, the weight gain was 3.87 mg/cm², and the surface hardness was 1027 HV0.05. Compared with nitriding without RE, the ε-Fe2,3N diffraction peak was enhanced in the RE nitriding layer. After 12 h of RE nitriding, La, LaFeO3, and a trace amount of Fe2O3 appeared. The corrosion rate of the modified layer was at its lowest (15.089 × 10⁻² mm/a), as was the current density (1.282 × 10⁻⁵ A/cm²); therefore, the corrosion resistance improved.

Keywords: 38CrMoAl steel; plasma nitriding; microhardness; corrosion resistance; modified layer

1. Introduction

Plasma nitriding is a widely used chemical heat treatment technology. This method uses the active nitrogen atoms generated during the cathode sputtering process at a relatively low temperature (350–570 °C) to accumulate on the surface of a workpiece, diffuse into the substrate, and finally form a nitriding-modified layer with excellent performance. For a variety of alloy materials, it can improve surface hardness, fatigue resistance, wear resistance, and corrosion resistance [1–7]. At present, the methods most commonly used to increase the thickness of the nitride layer are to increase the nitriding temperature or to extend the process time [8]. However, these methods have some disadvantages, such as the coarsening of the structure and reduced hardness [9,10]. Therefore, the question of how to increase the thickness of the modified layer in a short time presents a difficult challenge. Researchers have found that adding some catalysts can improve the efficiency of nitriding. Among them, rare earth (RE) elements have been proven to be effective catalysts for chemical heat treatment [11].

In the early 1980s, researchers studied the chemical heat treatment of RE. In the following decades, scholars explored the effects of RE catalysis and microalloying [12]. Many researchers observed that in the process of gas carburizing, gas carbonitriding, gas nitrocarburizing, and plasma nitriding, RE elements can diffuse deeper to the surface [13–15]. At the same time, the involvement of RE elements causes the nitrogen to diffuse deeper into the nitrided layer. Yan et al. [16,17] showed that rare earth elements induced nitriding and observed similar catalytic effects; they also studied the general catalytic effect of rare
earth elements on the nitriding process of various steels, including stainless steel [18,19] and alloy steel [20–22].

Due to its good wear resistance, high fatigue strength and high strength, 38CrMoAl steel is widely used in many industries [23,24]. However, in some applications, it cannot meet the requirements of high surface hardness and corrosion resistance. Therefore, surface modification approaches are used to improve the structure and properties of 38CrMoAl steel. However, there are few studies on the surface modification of 38CrMoAl steel, especially with the addition of RE. Therefore, it is very important to study the effect of plasma nitriding following the addition of RE on the thickness, surface hardness, phase composition, and corrosion resistance of a modified 38CrMoAl steel layer.

2. Materials and Methods

The test material was 38CrMoAl steel with a thickness of 5 mm and a diameter of φ 20 mm, and its chemical compositions are shown in Table 1. The steel was solution-treated in an SXL-1400C box-type electric furnace and austenitized at 940 °C for 1 h, after which oil quenching treatment was performed. A QG-1 metallographic cutting machine was used to cut the solution-treated sample into 5 mm thin slices. The surface was polished with 120# water sandpaper. Next, absolute ethanol was used for ultrasonic cleaning for 10–15 min. The sample was then blow dried with a hair dryer before being wiped with a dust-free cloth, weighed, and tied together with a thin iron wire. The 1 × 1 × 1 cm³ rare earth (RE) lanthanum was cut into 1/8 cube blocks, and the RE was hung on the cathode platform in order to maximize the sputtering of RE.

### Table 1. Chemical composition of 38CrMoAl steel (wt.%).

| Element | C  | Si  | Mn  | Cr  | Mo  | Al  | Fe  |
|---------|----|-----|-----|-----|-----|-----|-----|
| Content | 0.38 | 0.31 | 0.45 | 1.67 | 0.20 | 0.88 | Bal. |

Before plasma nitriding in the LDMC-30AFZ ion nitriding furnace, the furnace was evacuated to below 10 pa. The nitriding atmosphere was NH₃ (100 mL/min) at 550 °C for 4, 8, and 12 h. A CPA-225D electronic balance (Sai Dolis scientific instruments Beijing co., Ltd., Beijing, China) was used to measure the mass before and after the nitriding sample. After nitriding, the surface of the sample was polished with 120#-2000# water sandpaper. The polished sample was corroded with 4% nitric acid alcohol solution for 15 s, rinsed with alcohol, and dried. The cross-section structure and thickness of the modified layer was observed with a 9XB-PC metallographic microscope, the surface morphology was observed with a ThermoScientific™ Apreo C (TEM, JEOL Ltd, Tokyo, Japan) model field emission scanning electron microscope, and a HV-1000IS microhardness tester (Shanghai Jing Precision Instrument Manufacturing Co., Ltd., Shanghai, China) was used to test the hardness of the modified layer. The phase composition of the sample surface after nitrogen treatment was characterized by a D8 Advance X-ray diffractometer (Bruker, Karlsruhe, Germany). Anodic polarization tests were employed to estimate the corrosion resistance of the specimens in a 3.5 wt.% NaCl solution. The reference electrode was Ag/AgCl and a platinum column was used as the auxiliary electrode. A corrosion resistance test was performed in a CHI604E B16276 electrochemical workstation.

3. Results and Discussion

3.1. Microstructure and Phase Composition of Modified Layer

Figure 1 shows the metallographic microstructure of the modified layer of 38CrMoAl steel without rare earth (RE) and with rare earth. The nitrided layer without RE was clearly divided from the matrix, and the microstructure of the nitrided layer was uniform. After nitriding for 4, 8, and 12 h, the thickness of the modified layer was 222.2, 345.7, and 355.9 μm, respectively; the thickness of the modified layer increased gradually with the extension of nitride time. However, at the same temperature of 550 °C, after the addition
of RE, the nitrided layer thickened with time to 252.4, 358.0, and 391.8 µm (4, 8, 12 h), respectively. The thickened histograms of the nitrided modified layer with RE and without RE after nitriding for 4, 8, and 12 h are shown in Figure 2. The modified layer was thickened by 30.2, 12.3, and 34.9 µm at 4, 8, and 12 h, respectively. Compared with the nitriding layer without the addition of RE, the thickening rates of the RE nitriding layer were 13.6%, 3.50% and 9.80%, respectively, indicating that it was easier to obtain a thicker nitriding layer with the addition of RE at the same nitriding time intervals. The addition of RE in nitrided steel increased the concentration of microscopic crystal defects, such as vacancies and dislocations, in the modified layer and caused lattice distortion of the iron atoms around it, which was beneficial for the adsorption and diffusion of the nitrogen atoms. Therefore, in a short period of time, a higher nitrogen concentration was formed in the matrix, providing a very high nitrogen potential and concentration gradient for the inward diffusion of nitrogen atoms, which also provided good thermodynamic conditions for the diffusion of nitrogen atoms.

As shown in the XRD patterns of Figure 1a–c, which correspond to Figure 3a without RE nitriding, the main phase of the modified layer without RE nitridation was γ’-Fe₄N and the secondary phase was ε-Fe₂₃N. However, La and LaFeO₃ oxides were detected in the XRD patterns shown in Figure 1d–f, corresponding to Figure 3b with the addition of RE nitridation for 12 h. Because LaFeO₃ showed a strong adsorption effect on N atoms [25], the thickening rate of the modified layer increased with the addition of RE nitriding within the same timeframe.

**Figure 1.** Microstructure of modified layer of 38CrMoAl steel under different nitriding conditions: (a) PN, 4 h; (b) PN, 8 h; (c) PN, 12 h; (d) RE, 4 h; (e) RE, 8 h; (f) RE, 12 h.
Figure 2. Cross-section thickening histogram of nitrided layer of 38CrMoAl Steel without and with RE.

Figure 3. XRD pattern of 38CrMoAl steel without and with RE nitriding. (a) Without RE (b) With RE.

3.2. Weight Gain and Microhardness of Modified Layer

Figure 4 shows the histogram (4, 8, and 12 h) of the thickening of the steel without the RE-nitridated layer. The weight gain was 2.42, 3.35 and 3.75 mg/cm$^2$ at each respective time interval. Figure 5 shows the microhardness diagram of the modified layer. After 4 h of nitriding, the surface hardness was 701.3 HV$_{0.05}$. After nitriding for 8 h, the surface hardness was 975.3 HV$_{0.05}$. After 12 h of nitriding, the surface hardness of the modified layer was 882.5 HV$_{0.05}$. 
Figure 4. Histogram of weight gain of modified layer under nitriding conditions without RE.

Figure 5. Microhardness of modified layer cross-section under nitriding conditions without RE.

Figure 6 shows the histogram of the weight gain after nitriding with the addition of for 4, 8, and 12 h. The weight gain was 2.49, 3.43, and 3.87 mg/cm$^2$, respectively. Figure 7 shows the microhardness distribution under nitriding conditions with RE. After nitriding for 4, 8, and 12 h, the surface hardness was 840.9, 989.6, and 1027.0 HV$_{0.05}$, respectively, and the surface hardness reached its peak at 12 h. According to the microhardness bar chart of the nitriding surface without and with rare earth in Figure 8, the increase percentage of the surface hardness after nitriding with RE over time was 19.7%, 1.47%, and 16.3%, for each respective time interval. Therefore, the surface hardness improved after the addition of RE nitriding. This was mainly because the addition of RE refines the microstructure of the nitrided layer and adjusts the phase proportion [26]. Moreover, there are dispersion strengthening and solid solution strengthening effects after adding rare earth nitriding. At the same time, the infiltration of rare earth elements leads to serious lattice distortion and increases dislocation around the boundary, resulting in grain boundary strengthening.
Figure 6. Bar graph of weight gain of modified layer under nitriding conditions with RE.

Figure 7. Microhardness of modified layer cross-section under nitriding conditions with RE.

Figure 8. Histogram of the surface microhardness of the modified layer under nitriding conditions with and without RE.
3.3. Element Distribution in Modified Layer and EDS Energy Spectrum Analysis

The modified surface layers of the 12 h samples without RE nitriding were magnified by 100. Point scanning and plane scanning were performed for a, b, c, d, and e in Figure 9a by an EDS analyzer. Figure 9b1–b5 and Figure 9c1–c4 show the modified layer EDS spectrum test results. The distance from the surface of the modified layer was gradually transferred to the matrix, and the content distribution of each element was uneven. In combination with the line scan and plane scan in Figure 9a and Figure 9b1–b5, it is also shown that under nitriding conditions without RE, the content of the element N in the modified layer did not change significantly, and the distribution of the elements C, Fe, and Cr in the modified layer was relatively uniform.

![Figure 9. Element distribution diagram of the 12 h-nitrided layer without the addition of RE.](image)

Figure 9 shows the surface morphology and elements content after 12 h of RE nitriding. As can be seen from Figure 10b1–b5, the La content in the modified layer was small, while the N content showed a downward trend. At position c in Figure 10a, dot analysis was performed at 130.60 µm of the permeable layer, and the content of La was consistent with the surface content of the nitrided layer, indicating that RE atoms can be diffused into the modified layer. This was confirmed by mid-plane scanning, shown in Figure 10d5, but with the thickening of the nitrided layer, the modified layer contained a trace of La. Comparison between Figures 9b1 and 10b1 shows that the content of N element on the surface without RE nitriding was higher than it was after RE nitriding, although the thickness of the modified layer on the surface without RE nitriding was thinner than
that of the layer subjected to nitriding with RE [27]. This indicates that the increase of the nitrogen concentration or nitrogen the potential did not necessarily lead to the thickening of the modified layer. The thickness of the modified layer is not only related to the external N potential, but also to the internal diffusion of N atoms [28].

![Figure 10. Element distribution diagram in 12 h-nitrided layer with the addition of RE.](image)

(a) Fe, N, line-scan; (b1-b5) a,b,c,d,e points correspond to dot energy spectrum; (c) The distance between La and the surface of the modified layer; (d1-d5) C, N, Cr, Fe, La element surface scanning.

The surface structure of the 12 h RE-nitrided samples was further studied by transmission electron microscopy (TEM). The bright-field image and the corresponding selected region electron diffraction (SAED) map are shown in Figure 11. A large number of nanocrystals were observed, as shown in Figure 11a. The size of the precipitated phase was extremely small, with the maximum size no greater than 60 nm. The results showed
that the addition of RE refined the grain size and produced a microalloying effect on the modified layer. Figure 12b shows a clear diffraction ring caused by a large number of nanocrystals, indicating that these La, γ′-Fe4N, ε-Fe23N, and La phases existed in the outermost layer in the form of nanocrystals. The size of these phases was generally limited to 100 nm, so they can be referred to as nanophases. Yan et al. [29,30] obtained similar results with plasma nitriding, and found that the existence of the nano-phase helps to improve the hardness and wear resistance of the nitriding layer. Furthermore, these phases are consistent with the XRD results in Figure 3b.

**Figure 11.** TEM image after rare earth nitriding for 12h. (a) Bright-field image; (b) Corresponding SAED.

**Figure 12.** Corrosion resistance curve of modified layer under different nitriding conditions (4, 8, 12 h): (a) PN; (b) RE.

### 3.4. Polarization Curve and Fitting Data

The polarization curve data in Figure 12a were used for fitting, and the results are shown in Table 2. After nitriding for 4, 8, and 12 h, the corrosion rates of the modified layer were \(21.405 \times 10^{-2}\) mm/a, \(21.939 \times 10^{-2}\) mm/a, and \(25.714 \times 10^{-2}\) mm/a, respectively. The corrosion rates peaked after 12 h of nitriding. The current corrosion density was also increased to \(2.186 \times 10^{-5}\) A/cm² compared with that of the non-nitried sample. After fitting the polarization curve data shown in Figure 12b, we obtained the fitting results shown in Table 3. After nitriding with RE for 4, 8, and 12 h, the corrosion rates of the modified layer were \(16.092 \times 10^{-2}\) mm/a, \(20.754 \times 10^{-2}\) mm/a, and \(15.089 \times 10^{-2}\) mm/a,
respectively; the corrosion rate decreased, increased and then decreased again. The corrosion rates of the RE-nitrided samples reached their minimum at 12 h, and the current density of the RE-nitrided samples decreased to $1.282 \times 10^{-5}$ A/cm² compared with the samples without RE, and the width of passivation zone was wider. Tang [25] discussed the main reasons for the enhanced corrosion resistance of ε-nitride compounds on the workpiece surface. The X-ray diffraction analysis in Figure 3b shows that the presence of La and oxide LaFeO₃ significantly improved the intensity of the ε-Fe₂₋₃N diffraction peak and increased the content of the ε-phase in the samples. Because ε-nitrides are the most important parameters controlling the corrosion resistance of nitrided samples, the corrosion resistance of our rare earth-nitrided samples was enhanced at 12 h.

| Polarization Curve Fitting Data Unit | Untreated | 4 h | 8 h | 12 h |
|-------------------------------------|-----------|-----|-----|------|
| Corrosion rate ($\times 10^{-2}$ mm/a) | 17.472    | 21.405 | 21.939 | 25.714 |
| Rp($\times 10^3$ Ω/cm²)             | 1.756     | 1.434 | 1.400 | 1.193 |
| Io($\times 10^{-5}$ A/cm²)           | 1.486     | 1.820 | 1.865 | 2.186 |
| Eo/V                                 | −0.674    | −0.960 | −0.953 | −0.967 |
| Passivation zone width/V             | 0.715     | 2.219 | 2.267 | 2.395 |

| Polarization Curve Fitting Data Unit | Untreated | 4 h | 8 h | 12 h |
|-------------------------------------|-----------|-----|-----|------|
| Corrosion rate ($\times 10^{-2}$ mm/a) | 17.472    | 16.092 | 20.754 | 15.089 |
| Rp($\times 10^3$ Ω/cm²)             | 1.756     | 1.906 | 1.479 | 2.034 |
| Io($\times 10^{-5}$ A/cm²)           | 1.486     | 1.368 | 1.765 | 1.282 |
| Eo/V                                 | −0.674    | −0.953 | −0.954 | −0.962 |
| Passivation zone width/V             | 0.715     | 2.955 | 2.934 | 2.458 |

3.5. Thermodynamics of the Modified Layer

Because the nitrided atmosphere was NH₃, the main reaction in the furnace was:

$$\text{NH}_3 \rightarrow 3/2 \text{H}_2 + [\text{N}] \quad (1)$$

The decomposition of ammonia produced [N] atoms (Equation (1)), some of which penetrated into the surface of the workpiece, and the other part formed N₂, which became a stable gas.

The nitride produced by the decomposition of iron nitride provided the conditions for the formation of rare earth nitride [31].

$$2 \varepsilon\text{-Fe}_2\text{N} \rightarrow \text{Fe}_4\text{N} + 2[\text{N}] \quad (2)$$

$$\gamma'\text{-Fe}_4\text{N} \rightarrow 4\alpha\text{-Fe} + [\text{N}] \quad (3)$$

The RE formed compounds with nitrogen (N), a gap element of chemical heat treatment, and the bond changed from metal to ionic. Rare earths also reacted with hydrogen (H) to form rare earth hydrides.

Since the hydrogenation of rare earth can occur above 300 °C:

$$\text{La}^{3+} + 2\text{Cl}^- + \text{H}_2 = \text{LaH}_{2.3} + 3/2\text{Cl}_2 \quad (4)$$

Trace RE hydrides floating in the furnace flow improved the adsorption capacity of the workpiece surface diffusion. When RE hydrides are attracted to the workpiece surface, the following rare earth nitriding reactions occur:

$$\text{LaH}_2 + [\text{N}] \rightarrow \text{La}[\text{N}] + \text{H}_2 \quad (5)$$
The thermodynamic driving force of the decomposition of $\varepsilon$-Fe$_2$N (Reaction 2) was greater than that of $\gamma'$-Fe$_4$N (Reaction 3). Therefore, it is speculated that $\gamma'$-Fe$_4$N was still in the modified layer [32]. The calculated negative $\Delta G^0$ shows that the reaction occurs spontaneously from left to right in the range of 575–700 °C, forming active compounds of rare earth and nitrogen.

4. Conclusions

We studied the effect of the addition of rare earth (RE) to a modified layer of 38CrMoAl steel by plasma nitriding at 550 °C for 4, 8, and 12 h, and in an HN$_3$ atmosphere. The following conclusions were drawn:

(1) For nitriding without the addition of RE for 4, 8, and 12 h, the thickness of the modified layers was 222.2, 345.7, 355.9 µm for each respective timeframe, the weight gains were 2.42, 3.35, and 3.75 mg/cm$^2$, respectively, and the maximum surface hardness was 882.5 HV$_{0.05}$, at 12 h. After nitriding with the addition of RE for 4, 8, and 12 h, the modified layers were thickened by 30.2, 12.3 and 34.9 µm, respectively, and the weights were 2.49, 3.43 and 3.87 mg/cm$^2$, respectively. The maximum surface hardness was 1027.0 HV$_{0.05}$, at 12 h.

(2) After 12 h of nitriding with RE, La, LaFeO$_3$, and trace Fe$_2$O$_3$ appeared in the modified layer. The diffraction peak of $\gamma'$-Fe$_4$N in the RE nitriding layer was weakened compared with the layer nitrided without RE, while the diffraction peak of $\varepsilon$-Fe$_2$N was enhanced. Meanwhile, La, $\gamma'$-Fe$_4$N, $\varepsilon$-Fe$_2$N, and La phases existed in the outermost layer in the form of nanocrystals.

(3) After nitriding with the addition of RE for 12 h, the corrosion rate of the modified layer decreased to a minimum of $15.089 \times 10^{-2}$ mm/a, and the current density also decreased to a minimum of $1.282 \times 10^{-5}$ A/cm$^2$; consequently, the corrosion resistance increased.

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