The role of chromium content in the long-term atmospheric corrosion process

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The corrosion of alloy steels with different amounts of Cr was studied using electrochemical tests, wet–dry cycle corrosion, X-ray diffraction, and Kelvin probe force microscopy. The results show that the content of Cr is positively correlated with the corrosion resistance of steel, but the corrosion resistance of atmospheric corrosion does not show the same pattern. The atmospheric corrosion resistance of Cr-containing steel exhibits three different stages with the change of Cr content. When the Cr content is in the range of 1–4%, the corrosion rate is high and does not change within the Cr content. As the Cr content was further increased from 4 to 7%, the corrosion rate exhibited a linear decrease and then drops rapidly when the Cr content reaches 8%. These three different corrosion resistance stages are related to the influence of Cr content on Fe2O3 content in the rust layer.

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INTRODUCTION

It is well known that the corrosion resistance of materials can be improved by adding corrosion-resistant alloying elements1–3. In recent years, a method to improve the corrosion resistance of materials is to form a dense and stable rust layer. One of the typical examples is weathering steel. By adding certain amounts of corrosion-resistant elements, a dense rust layer is formed on the steel surface, which can impede further corrosion processes of the substrate to achieve improvement of the steel corrosion resistance4–8. The recent researches have suggested that addition of Cr, Cu, or P alloying elements can significantly improve the rust layer structure stability of the weathering steel, thereby improving its applicability in severe industrial and marine environments9–12.

Several studies have focused on the effect of alloying elements on the corrosion resistance. A study by Ma et al.16 on the atmospheric corrosion behavior of weathering steel in the tropical marine environment found that the weathering resistance of weathering steel is related to the deposition of chloride ions and the enrichment of Cr in the rust layer and that the latter can significantly improve the corrosion resistance of weathering steels in environments with low chloride ion deposition. Several studies17–21 found that Cu can enhance the stability of the rust layer when used together with Cr, thus improving the corrosion resistance of weathering steel. However, a minimum of 0.10 wt% Cu is required in order to significantly improve the corrosion resistance of the steel22. The rust layer of corrosion-resistant steel after long-term exposure to the atmosphere has a two-layer construction, and a significant difference Cr content presents in the inner and outer layers, respectively23. A Cr-enriched inner layer mainly consists of α-(Fe1–xCrx)OOH, a Cr substituted FeOOH, which acts as a final stable protective layer. Other studies also found that a pseudo-passivation film formed on some Cr-containing steels in the CO2 corrosion environment, and this film is consisted of Cr(OH)324,25. After analyzing the rust layer structure of Cr-containing steel, Jiang et al.26 found that Cr is mainly in the forms of Cr(OH)3, CrOOH, or Fe2Cr2O5 in the rust layer, that is, Cr mainly exists in the form of Cr7+ among the participants during the iron oxide process. Yamashita et al.27 believed that the existence of Cr element can replace the nano-sized goethite particles to create a denser protective rust layer in the inner rust layer. Although all the existing studies so far used several alloying elements together28,29, it is not difficult to see from these studies that the Cr content and the corrosion resistance have a close correlation. However, the study of the influence of chromium on the initial stages of the atmospheric corrosion of weathering steels by Kamimura and Stratmann30 found that the positive effect of chromium is not only not apparent in saline environments but also seems to be even harmful by accelerating weathering steels corrosion in coastal environments. Other researchers31–32 also pointed out that the presence of 0.60–1.30 wt% Cr was sometimes slightly detrimental, especially in industrial atmospheres. Therefore, the effect of the Cr content on the formation of the weather-resistant rust layer in steel, as well as the relationship between the Cr content and the density of the rust layer, has not yet reached a consentient understanding.

In this work, the steel samples with eight different Cr contents were studied to identify the relationship between the Cr content and the atmospheric corrosion resistance. A dry–wet cyclic corrosion test was used to study the effect of the Cr content on the corrosion resistance of the corrosion layer during the atmospheric corrosion process. The purpose of this study is to provide an insight into the influence of the Cr content on atmospheric corrosion resistance. In addition, this study may provide useful data for the development of micro-alloyed corrosion-resistant materials.

RESULTS

Microstructure

The metallographic structure of the eight experimental materials is shown in Fig. 1. It can be seen from the figure that Cr changes the microstructure of the matrix. The Cr content primarily affects the proportion of bainite in the steel. The W1 structure is essentially ferrite with a very small amount of lath bainite. W2, W3, and W4 are both granular bainite and ferrite. As the Cr content increases, the proportion of granular bainite increases gradually.

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This occurs because the addition of Cr to a steel can delay the pearlite transformation and promote the bainite transformation, making it easier to form the bainite structure at a slow cooling rate after austenitizing. When the Cr content reaches $\geq 5\%$, the structure is all bainite, and the size of bainite becomes finer with the increase of Cr content; the W8 has the smallest grain size.

Wet-dry cycles corrosion results
To study the atmospheric corrosion behavior of the steels with different Cr contents, dry and wet cycle experiments were carried out on the eight steel samples. Figure 2 shows that, as the corrosion cycles increased, the corrosion rate gradually stabilizes. At the initial stage of the experiment, the main reaction is the

Fig. 1  Metallographic structure of samples with various Cr content. a–h correspond to W1–W8 samples, respectively, and as the Cr content increases, the proportion of granular bainite increases gradually.

Fig. 2  Corrosion rate of eight experimental steels during alternating wet–dry test in 0.5% NaCl ($\text{pH} = 3$) solution. a 24 h, b 72 h, c 120 h, d 168 h. The dotted line corrosion rate in b is 3.8 mm y$^{-1}$, the dotted line in c for 3.7 mm y$^{-1}$, and the dotted lines in d for 5 and 2.5 mm y$^{-1}$, respectively. Based on corrosion rate variation, there are two corrosion stages in b, c and three in d.
corrosion of the steel substrate. Therefore, the corrosion rate gradually decreases with the increase of Cr content. The Cr content showed a certain positive effect on the corrosion resistance in the early stage of corrosion. However, after the 72-h dry–wet cycle accelerated experiment, the corrosion rate change started to show two different trends for the samples with different Cr contents. The effect of Cr on corrosion rate of the samples with low Cr contents (W1–W4) decreased gradually and tended to be the same, while the effect of Cr on corrosion resistance in those with high Cr content (W5–W8) still showed a positive correlation with the Cr content. This is mainly due to the gradual formation of the rust layer on the steel surface during the prolonged experiment time, where the main factor that affects the corrosion resistance has shifted from the corrosion resistance of the steel substrate to the density of the rust layer. The results after 120-h experiment show no more significant difference in the corrosion rate when the Cr content is <4%. The result from long-period cycling corrosion test indicated that 4% Cr content appeared to be a threshold for its role in affecting the corrosion rate. After the 168-h dry and wet cycle corrosion experiment (Fig. 2d), a stable rust layer structure has been formed on the surface of the material, and the corrosion rate at this time can be divided into 3 groups: (1) The corrosion rates of W1–W4 are basically the same, at ~5.5 mm y\(^{-1}\). (2) The corrosion resistance of W4–W7 gradually increases, and the corrosion rate decreases linearly from 5.5 to 2.7 mm y\(^{-1}\); (3) The corrosion rate of W8 is only 0.3 mm y\(^{-1}\), which is merely one-eighth of W1.

The macroscopic morphology (Supplementary Fig. 1) of the eight experimental materials after wet–dry cycles corrosion test shows that, at 24 h, the surface of the experimental steel features a thin layer of corrosion products, which is relatively flat and does not bubble. The surface rust layers of W1–W6 are predominantly dark brown, with some yellow-brown rust spots; W7 has a small amount of brick red rust on the surface, and W8 exhibits only a very slight corrosion and the substrate is visible. At 72 h, the presence of corrosion products is clearly more noticeable compared with 24-h samples but still not dense. There is obvious surface bubbling for W1–W4. The surface rust layers for W5–W7 are relatively flat. W8 is evenly covered with yellowish brown rust and is very dense. After 120 h, bubbling was observed for W5–W7, and there was no change in the surface appearance for W8. After 168 h, the degree of corrosion had obviously increased, the rust layer morphology for W1–W4 exhibited no difference, the corrosion extent for W5–W7 gradually reduced, and for W8 the surface was still covered in the yellow-brown rust layer, and the corrosion was trivial. Furthermore, the microscopic morphology of corrosion products after 168 h (Supplementary Fig. 2) show that the surfaces of W1–W7 feature more corrosion products, significant surface bubbling, and many cracks. The surface of W8, however, is covered by fine and uniform corrosion products, which is very dense with no cracks, thus protecting the substrate. Figure 3 shows the microscopic morphologies of the experimental steels after the removal of the rust. The surfaces of W1–W4 exhibit many pitting pits and severe uniform corrosion morphology. W5–W7 present many localized corrosion features. There are virtually no obvious corrosion marks on the surface of W8, but a small number of small pits. These results show that, in the Cr content range from 1 to 4%, the uniform corrosion decreases and the local corrosion increases as the Cr content increases. In the Cr content range from 5 to 7%, the depth and the number of pitting pits decrease with increasing Cr content. When the Cr content reaches 8%, the corrosion behavior of the surface changes suddenly, resulting in improved corrosion resistance and less surface corrosion. These results are consistent with the other studies, which had a conclusion that the local corrosion of steel becomes more obvious, and the interface of the base rust layer becomes uneven after adding Cr to steel\(^{19}\). The scanning electron microscopic (SEM) observation results of the rust layer cross-section are shown in Fig. 4, and energy-dispersive spectroscopic (EDS) line scan analysis was performed on the marked areas in Fig. 4. It can be seen from the cross-sectional topography that there is no obvious delamination in the rust layer. For W1–W4, the number of the pitting pits increases gradually as the Cr content increases, the rust layer cross-section is uneven, and there are many microscopic cracks and holes. Samples W5–W7 exhibit significant local corrosion, the bonding between the rust layer and the substrate is weak, and large cracks exist in the rust layer. The rust layer of W8 is the thinnest, with a thickness only ~20 μm. The rust layer structure is very dense and exhibits good bonding with the matrix. It can be seen from the EDS line scan that the rust layer near the substrate is enriched in Cr. Cl\(^{-}\) exists in the entire rust layer and is slightly higher in the inner layer than that in the outer rust layer. This indicates that the rust layer is an ion conduction layer, and Cl\(^{-}\) can penetrate through the rust layer to the metal interface, causing corrosion reaction therewith; this gradually increases the density of the rust layer. Cracks and holes also appear in the cross-section of the rust layer. This is because the rust layer contains various corrosion products, and the densities of different corrosion products are different and smaller than that of the matrix. During the continuous accumulation of corrosion products on the surface of the steel substrate, stress is generated at the interface between the rust layer and the substrate, and corrosion products themselves have poor ductility, resulting in cracks and holes.

The X-ray diffraction (XRD) analysis on the rust layers of the samples after the wet–dry cycles corrosion test was performed, and the results are shown in Fig. 5. It can be seen from figure that, for all the samples analyzed, the rust layers contain α-FeOOH, γ-FeOOH, and Fe\(_3\)O\(_4\). However, the content of each substance is significantly different in the different samples. Therefore, the K value method was used in order to semi-quantitatively analyze the rust layers to calculate the proportion of different phases in the rust layers. The results are shown in Table 1.

It can be seen from the table that, during the initial 24 h of corrosion, the rust layers of W1–W6 contain a large amount of Fe\(_3\)O\(_4\), while those of W7 and W8 only contain α-FeOOH. At 72 and 120 h, surface corrosion of the samples became more extensive, and the proportion of Fe\(_3\)O\(_4\) is detectably higher, which is consistent with the increase in the black rust amount, as evidenced by the macroscopic morphology results (Supplementary Fig. 1). After 168 h, the proportion of Fe\(_3\)O\(_4\) becomes relatively small. Most of the corrosion products for W1–W7 are Fe\(_3\)O\(_4\), while most of the corrosion products for W8 are α-FeOOH and γ-FeOOH with almost no Fe\(_3\)O\(_4\). The proportion of α-FeOOH increases with the increase of number of corrosion cycles.

The XRD results show that increasing the Cr content has little effect on the γ-FeOOH content in the rust layer, which remains at a low level. Therefore, for each experimental steel we analyzed the relationship between the corrosion rate and the ratio of Fe\(_3\)O\(_4\) in the rust layer for different corrosion cycles (Fig. 6). For each corrosion cycle, the ratio of Fe\(_3\)O\(_4\) in the corrosion product of W8 is consistent with its corrosion rate. After 168 h, it can be seen clearly that the Fe\(_3\)O\(_4\) ratio can be categorized into three groups based on the Cr content. For W1–W4, the ratio of Fe\(_3\)O\(_4\) remains stable at ~92%. For W4–W7, the ratio of Fe\(_3\)O\(_4\) decreases linearly from 92 to 73%, while the rust layer of W8 contains only 0.9% Fe\(_3\)O\(_4\). This change of Fe\(_3\)O\(_4\) content in the rust layer is consistent with the results from the corrosion rate change curve.

DISCUSSION

The polarization curves (Supplementary Fig. 3) of the eight experimental materials in the 3.5% NaCl solution shown that the corrosion resistance of the steel matrix improves as the Cr content increases. This is consistent with the traditional understanding of the role of Cr element but not with the results observed in the
atmospheric corrosion process. In order to further investigate this difference, the microscopic structure change of the substrate caused by Cr was studied to further clarify the reason for improving the corrosion resistance of the substrate. Many studies investigated the possibility that the steel structure affects the corrosion resistance, and the addition of alloying elements changes the microstructure of the steel matrix, which in turn can affect its corrosion resistance. Our work showed that, when Cr was added as an alloying element to low-carbon steel, it delayed the pearlite transformation and promoted the bainite transformation, allowing the bainite structure to form more easily after austenitization. It can also be seen from Fig. 1 that the addition of Cr does changes the original phase structure of the material. When the Cr content is <4%, the material was composed of two phases of ferrite and bainite. When the Cr content was increased to 8%, the grain size further improved the corrosion resistance. Therefore, it is considered that the difference of the corrosion resistance in the experiment might be mainly contributed by the structural changes caused by the addition of different amount of Cr. To validate the hypothesis, W3 sample was selected for Kelvin probe force microscopy (KPFM) analysis. (Supplementary Fig. 4 and Fig. 7).

Supplementary Fig. 4 shows the metallographic structure of the W3 steel containing 3% Cr. The structure is composed of ferrite and granular bainite. The locus marked in the figure was where the KPFM testing performed. The KPFM result is shown in Fig. 7. Figure 7a–d are the height map, the potential map, the three-dimensional map of the potential, and the potential change curve.
of the section, respectively. It can be seen from Fig. 7d that the potential of the granular bainite is much higher than the potential of the matrix ferrite, and the difference between the two is $\sim 80$ mV. The existence of this potential difference may have two effects. (1) The corrosion resistance of the steel matrix improves with more phase content that has better corrosion resistance, as shown in Fig. 2a and Supplementary Fig. 3 that display an early corrosion rate decrease. The corrosion resistance of the steel matrix continues to improve with the addition of more Cr content$^{29,37 – 39}$. (2) The galvanic difference between the two phases caused the variations of the morphology and structure during the atmospheric corrosion process. Due to the difference of the potential between the two phases, the corrosion process is promoted and the corrosion morphology changes from uniform corrosion to local pitting, as shown in Fig. 3. At the later stage after a long-period dry–wet alternate corrosion process, the major factor that influence the corrosion rate became the structure of the rust layer. Therefore, a detailed analysis of the changes in the structure of the rust layer is also required.

The rust layer structure plays an important role in the corrosion resistance of materials$^{30,41}$. In atmospheric corrosion, the main process on the material surface is the oxidation of Fe and the transformation of the rust layer. In Evans model$^{42}$, it is considered that a carbon steel undergoes strong cathodic depolarization in the rust layer at the beginning of corrosion and then an anodic reaction in the rust layer. This process occurs at the metal/$\text{Fe}_3\text{O}_4$ interface:

$$\text{Fe} \rightarrow \text{Fe}^{2+} + 2e^-$$  \hspace{1cm} (1)

The cathodic reaction occurs at the $\text{Fe}_3\text{O}_4$/FeOOH interface:

$$\text{Fe}^{2+} + 8\text{FeOOH} + 2e^- \rightarrow 3\text{Fe}_3\text{O}_4 + 4\text{H}_2\text{O}$$  \hspace{1cm} (2)

$$3\text{Fe}_3\text{O}_4 + 3/4\text{O}_2 + 9/2\text{H}_2\text{O} \rightarrow 9\text{FeOOH}$$  \hspace{1cm} (3)
It is considered that α-FeOOH is an insulating inactive substance, which is the most stable hydroxy iron oxide and a main constituent of the protective rust layer. Tanaka et al.43 identified that Fe₂O₃ plays an important role in reducing the corrosion resistance of rust layers, while α-FeOOH and γ-FeOOH promote the formation of Fe₂O₄. Fe₂O₄ serves as a conductor in the rust layer. Its conductivity is similar to that of pure metal. The electrons in the metal can be transferred by Fe₂O₄ as carriers. Therefore, when the rust layer contains a large amount of Fe₂O₄, it does not provide a protective effect. Instead, Fe₂O₄ acts as an active point on the surface and provides carriers for the electrochemical process, thus promoting the corrosion process of metals. Addition of Cr can have two effects on the alteration of the electrochemical process, thus promoting the corrosion process active point on the surface and provides carriers for the corrosion resistance of rust layers, while Fe₂O₄→FeOOH and promoting the formation of FeOOH in the rust layer, and this conversion and Cr content present positive correlation.

METHODS

Materials preparation
Eight groups of experimental steel samples with different Cr contents marked as W1, W2, W3, W4, W5, W6, W7, and W8 were prepared. The compositions of these samples are shown in Table 2. They were prepared using vacuum melting and then casting into round ingots, followed by 1-h homogenization at 1200 °C. The steel ingots were rolled into 6-mm-thick strips with a final rolling temperature of approximately 900 °C, followed by water quenching to 520–540 °C and then air cooled to room temperature.

In order to eliminate internal stress from the rolling process, the samples were normalized at 850 °C for 0.5 h. The detailed process is described in Supplementary Fig. 5. The final samples were cut into 10 mm × 10 mm × 3 mm. All the samples were sanded to 2000#, polished, and etched with 4% nitric acid alcohol solution to observe the microstructure.

Wet–dry cyclic corrosion test
The dry–wet cycle experiment is generally considered as one of the more efficient methods for simulating atmospheric corrosion because of its similarity with the actual outdoor corrosion45,46. Each cycle of the wet/dry corrosion test is 1 h, which consists of two stages: (1) immersing the specimens in a 0.5% NaCl solution with pH = 3.0 at 40 °C for 45 min and (2) drying the specimens with four heating lamps for 15 min. The relative humidity inside the testing container was maintained at 70% (±5%) during the experiment. The samples have a size of 50 mm × 25 mm × 3 mm, and each experimental group was made of three samples. Prior to the experiments, all of the samples were processed as follows: sanding (to 1500#) → pure water cleaning → acetone degreasing → anhydrous ethanol dehydration → 24-h drying. The experimental periods were 24, 72, 120, and 168 h. For each sample, the initial weight was recorded as W₀ (accurate to 0.0001 g), and the sample size was measured using a Vernier caliper. After the experiment, all the samples were placed in an ultrasonic cleaner with a solution of 500 mL HCl → 500 mL H₂O₂ → 3.5 g hexamethylene-tramine at 25 °C for about 15 min to remove the corrosion product47. Then the samples were rinsed using deionized water, followed by anhydrous ethanol alcohol and dried in cold air. The final masses Wᵣ were recorded (accurate to 0.0001 g). Meanwhile, an untreated sample also went through the same processes for each steel to calibrate the mass loss produced during the process of removing corrosion products. The corrosion rates (R, mm y⁻¹) were calculated as follows:

$$ R = \frac{8.76 \times 10^4 W₀ - Wᵣ}{Spt} $$

where (W₀ − Wᵣ) represents the weight loss of the sample after the dry–wet cyclic corrosion test, measured in grams; S represents the exposed area of the sample and is in cm²; p represents the sample’s density and was taken as 7.8 g cm⁻³; and t is the test time in years.

Table 1. Semi-quantitative analysis of rust layer XRD results.

| Number | Period | Fe₂O₄ (wt.%) | α-FeOOH (wt.%) | γ-FeOOH (wt.%) |
|--------|--------|--------------|----------------|----------------|
| W1     | 24 h   | 84.57157     | 15.42843       | 0              |
| 72 h   | 94.97698 | 24.33409     | 5.53714        | 0              |
| 120 h  | 94.46285 | 0            | 3.83782        | 96.16466       |
| 168 h  | 92.74633 | 2.08922      | 5.164446       | 97.98532       |
| W2     | 24 h   | 90.46199     | 6.302565       | 3.235448       |
| 72 h   | 97.9693  | 0            | 5.164446       | 97.98532       |
| 120 h  | 95.57277 | 0            | 4.427234       | 96.00000       |
| 168 h  | 92.2527  | 3.897821     | 3.849475       | 94.95052       |
| W3     | 24 h   | 89.17726     | 10.82274       | 0              |
| 72 h   | 96.78247 | 0            | 3.217528       | 97.98532       |
| 120 h  | 96.3087  | 0            | 3.691301       | 96.30870       |
| 168 h  | 90.77812 | 4.26159      | 96.00000       | 96.00000       |
| W4     | 24 h   | 87.33965     | 6.547475       | 0              |
| 72 h   | 91.09919 | 2.112929     | 6.787885       | 97.98532       |
| 120 h  | 96.28528 | 0            | 3.717421       | 96.00000       |
| 168 h  | 94.20305 | 2.495278     | 3.301669       | 94.20305       |
| W5     | 24 h   | 77.43117     | 22.56883       | 0              |
| 72 h   | 92.53767 | 0            | 4.762329       | 97.98532       |
| 120 h  | 95.84923 | 0            | 4.150765       | 95.84923       |
| 168 h  | 88.43245 | 11.56755     | 96.00000       | 96.00000       |
| W6     | 24 h   | 76.53408     | 16.6905        | 0              |
| 72 h   | 94.01022 | 0            | 5.989777       | 97.98532       |
| 120 h  | 93.8505  | 0            | 6.149432       | 93.85050       |
| 168 h  | 76.78732 | 14.32297     | 8.889715       | 94.95052       |
| W7     | 24 h   | 73.21516     | 15.47285       | 0              |
| 72 h   | 92.93271 | 0            | 7.067288       | 97.98532       |
| 120 h  | 95.23767 | 0            | 4.762329       | 95.23767       |
| 168 h  | 76.78732 | 14.32297     | 8.889715       | 94.95052       |
| W8     | 24 h   | 73.77694     | 14.75088       | 0              |
| 72 h   | 92.2527  | 0            | 11.47218       | 94.95052       |
| 120 h  | 94.01022 | 0            | 4.762329       | 94.01022       |
| 168 h  | 73.77694 | 14.75088     | 8.889715       | 94.95052       |
Fig. 6  Variation tendency of Fe₃O₄ content in corrosion products and corrosion rate for eight experimental steels during various wet–dry corrosion cycles. a 24 h, b 72 h, c 120 h, d 168 h. The orange lines in a–d all represent the change curve of corrosion rate, and the black line represents the change of Fe₃O₄ content of corrosion products.

Fig. 7  The KPFM of different structure in 3% Cr content experimental steel. a Height map, b Potential map, c 3D map of potential, d potential change curve of the red section line in b.
Corrosion morphology and rust analysis
Surface and cross-sectional morphologies of the samples were observed using a KEYENCE VHX-200 stereomicroscope and a FEI Quanta250 SEM. Their rust layers were analyzed with an EDS. The phases in the corrosion products were determined by XRD on a Rigaku analyzer over a 10°–90° range with a scan step of 0.02°. The patterns were refined using the Rietveld refinement method.

Electrochemical test
Electrochemical tests were performed using a M2273 electrochemical workstation and a three-electrode system, where the samples were used as the working electrodes, a Pt plate as the counter electrode, and a saturated calomel electrode (SCE) as the reference electrode. The test solution was 3.5% NaCl (pH ∼7) containing 50 ppm dissolved oxygen and 30 ppm Fe ions. The reference electrode was a saturated calomel electrode (SCE) in artificial seawater. The test was conducted at a constant current density of 10 μA cm⁻². The open circuit potential (OCP) was monitored for at least 20 min until it reached a stable state. Potentiodynamic polarization measurements were performed at 0.5% NaCl (pH = 3) using a KEYENCE VX-200 stereomicroscope and a FEI Quanta250 SEM. Their corrosion behavior was studied using the potentiodynamic polarization method.

KPFM test
Atomic force microscope (Multimode 8, Bruker) is used in the KPFM measurements. The equipment was located in a clean room at a constant temperature of 25 °C and an ambient relative humidity of approximately 20%. The probes used in the KPFM measurements were PFQNE-AL with a force constant of 1.5 N m⁻¹ and a resonant frequency of 300 kHz. The dual-scan mode was used to record the second signal in addition to the surface topographic signal. In a KPFM measurement, when a potential difference exists between the tip and the sample, an AC voltage is applied to the tip to induce harmonic oscillation of the cantilever. Afterwards, a voltage of the same magnitude is applied to the tip to stop the oscillation and reduce the potential difference. Using this method, the work function of the sample can be derived. Some studies have shown that the work function is closely related to the corrosion potential. The lower the work function of a material, the lower the corrosion potential and the easier the corrosion.

DATA AVAILABILITY
The data of this study are available from the corresponding authors upon reasonable request.

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