The relationship between activation–passivation transition and grain boundary dissolution on four steel samples in acidic solutions containing NO₂⁻

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Herein, for four steels (L80, N80, X65 and Q235) in acidic solutions (HNO₃, HCl, HAc and CO₂) containing NO₂⁻, the relationship between the activation–passivation (A–P) transition and the grain boundary dissolution (GBD) was studied by potentiodynamic polarization curve (PPC) measurements and scanning electron microscopy (SEM) observations. In the specific pH range of acidic solutions, where the four steels showed an electrochemical characteristic of the A–P transition, GBD was observed on the steel surface; however, at low or high pH values of the acidic solutions, the four steels respectively showed the electrochemical behavior of activation (A) or self-passivation (sP), and GBD was not observed on the steel surface. The effects of the acid type, pH value and steel type on the electrochemical characteristic of the A–P transition and the occurrence of GBD were also discussed in detail. Via this study, it was confirmed that under the electrochemical characteristic of the A–P transition, the occurrence of GBD was a general corrosion behavior of carbon steels and alloy steels in acidic solutions containing NO₂⁻.

1. Introduction

Steel products are widely applied in the fields of production and living; however, the problems of corrosion and failure inevitably limit their application and development, which are due to the presence of aggressive species in service environments.

In the electrochemical process of steel corrosion, the main cathodic reaction is H⁺ reduction or O₂ reduction, which is very closely related to the solution pH value. In strongly acidic solutions (relatively low pH), the dominant cathodic reaction is the H⁺ reduction with the standard potential \( E_{\text{st}} \) of \(-0.244 \text{ V}_{\text{SCE}}\). \(^{(1)}\)

\[
2\text{H}^+ + 2e^{-} \rightarrow \text{H}_2 \quad (E_{\text{st}} = -0.244 \text{ V}_{\text{SCE}}) \tag{1}
\]

According to the Nernst equation, the equilibrium potential \( E_{\text{eq}} \) of H⁺ reduction can be described as follows: \(^{(1)}\)

\[
E_{\text{eq}/\text{SCE}} (\text{H}^+/\text{H}_2) = -0.244 - 0.059 \text{ pH} \tag{2}
\]

In contrast, in weakly acidic and alkaline solutions (relatively high pH), the O₂ reduction with the \( E_{\text{st}} \) of \(0.157 \text{ V}_{\text{SCE}}\) becomes the dominant cathodic reaction: \(^{(1)}\)

\[
\text{O}_2 + 2\text{H}_2\text{O} + 4e^{-} \rightarrow 4\text{OH}^- \quad (E_{\text{st}} = -0.157 \text{ V}_{\text{SCE}}) \tag{3}
\]

The \( E_{\text{eq}} \) of O₂ reduction can be depicted as follows: \(^{(1)}\)

\[
E_{\text{eq}/\text{SCE}} (\text{O}_2/\text{OH}^-) = 0.984 - 0.059 \text{ pH} \tag{4}
\]

It is speculated that the following anodic reaction of Fe oxidation is independent of the solution pH value:

\[
\text{Fe} \rightarrow \text{Fe}^{2+} + 2e^{-} \quad (E_{\text{st}} = -0.684 \text{ V}_{\text{SCE}}) \tag{5}
\]

With an increase in the pH value, the \( E_{\text{eq}} \) for both the H⁺ reduction (eqn (2)) and the O₂ reduction (eqn (4)) would decline; this would result in the negative shift of the corrosion potential \( (E_{\text{corr}}) \) in the polarization curve. In addition, the left shift of the chemical equilibrium for the H⁺ reduction (eqn (1)) and the O₂ reduction (eqn (3)) would occur, resulting in a decrease in the corrosion current density \( (i_{\text{corr}}) \) in the polarization curve. Related studies involving the abovementioned discussion have been repeatedly reported.\(^{(12-14)}\)

On the other hand, intergranular corrosion (IGC) is a common and important type of localized corrosion for metals and alloys, which is attributed to the establishment of an electrochemical micro-couple between grain interiors (GIs) and grain boundaries (GBs).\(^{(15)}\) At present, studies on IGC are mainly focused on stainless steels (SSs)\(^{(16-20)}\) and aluminum alloys (AAs);\(^{(21-23)}\) this is due to high IGC susceptibility of SSs and AAs derived from the precipitation of harmful phases along GBs during the process of inappropriate thermal treatment. For SSs, precipitates such as Cr₂₃C₆,\(^{(16)}\) CrN/Cr₂N,\(^{(17)}\) σ phase,\(^{(18)}\) φ phase,\(^{(19)}\) and π phase\(^{(20)}\) have been reported to result in IGC susceptibility. Moreover, the IGC susceptibility of AAs has been reported to...
result from the precipitation of the T phase,23 S phase,22 β phase,23 T phase,24 η phase25 and other similar phases; due to the presence of IGC susceptibility, the electrochemical microcouple between GIs and GBs can be spontaneously established when SSs and AAs are applied in corrosive environments, thus inducing the occurrence of IGC.26

Except for SSs, IGC susceptibility is not prominent in other types of steel, and relatively few studies have been reported on IGC. In a previous study, for the Q235 carbon steel in a CO2–NaNO2 solution, the occurrence of the grain boundary dissolution (GBD) was observed on the steel surface when the Q235 steel was polarized in the activation–passivation (A–P) region.27 Subsequent studies further confirmed the occurrence of GBD on the surface of carbon steels when this type of steel was polarized in the A–P region in acidic solutions containing NO2− such as a CO2–NaNO2 solution,28 HNO3–NaNO2 solution29 and HCl–NaNO2 solution.30 However, to date, it is not clear that in acidic solutions containing NO2−, whether the occurrence of GBD is a special corrosion behavior of carbon steels or a general corrosion behavior of other steels, particularly alloy steels. To clarify the abovementioned statement, in this study, three low alloy steels (L80, N80 and X65) were chosen to study the relationship between the electrochemical characteristic of the A–P transition and the occurrence of GBD in four acidic solutions (HNO3, HCl, HAc and CO2) containing NO2−. Electrochemical measurements using a potentiodynamic polarization curve (PPC) and microstructural observation using a scanning electron microscope (SEM) were carried out; in addition, to understand the difference between the A–P transition and GBD of carbon steel and alloy steel, the PPC measurement and the SEM observation for the Q235 steel were performed.

2. Experimental

The detailed chemical composition of the L80, N80, X65 and Q235 steels is listed in Table 1. Herein, four steel samples were manually abraded up to 1000 grit with SiC abrasive papers, rinsed with deionized water and degreased in acetone.

The tested solutions were four acidic solutions containing NO2−: HNO3–NaNO2, HCl–NaNO2, HAc–NaNO2 and CO2–NaNO2. For the first three solutions, the diluted HNO3, HCl and HAc solutions were introduced to a 0.01 M NaNO2 solution to adjust the pH value. For the CO2–NaNO2 solution, the CO2 gas was purged into a 0.01 M NaNO2 solution until the CO2 saturated condition was reached and the pH value of the CO2–NaNO2 solution was 3.7.31

The PPC tests were carried out using the CS310 electrochemical workstation. A typical three-electrode system was applied, and the system was composed of a saturated calomel electrode (SCE) as the reference electrode, a platinum sheet as the counter electrode and a steel sample as the working electrode. Before each PPC test, the working electrode was immersed in the corresponding tested solution for 30 min to ensure a stable condition of the open circuit potential (OCP). The potential scanning rate was 0.1 mV s−1, and the potential scanning range was from −0.3 V OCP to the potential value corresponding to the occurrence of transpassivation or pitting. All the PPC tests were performed at ambient temperature. After this, the surface morphologies of the four steel samples were observed by the LEO-1450 SEM instrument.

3. Results and discussion

Fig. 1 shows the PPCs of L80, N80, X65 and Q235 steels in HNO3–NaNO2, HCl–NaNO2 and HAc–NaNO2 solutions with 0.01 M NaNO2 and different pH values. The effects of the acid type, pH value and steel type on the electrochemical behaviors were investigated, and the following five electrochemical behaviors were demonstrated in the PPCs: activation–passivation–transpassivation (A–P–T), self-passivation–transpassivation (sP–T), activation (A), activation–passivation–pitting (A−P–P), and self-passivation–pitting (sP–P). The detailed results of the electrochemical behavior are presented in Table 2.

According to previous studies,27–30 for carbon steels in acidic solutions containing NO2−, the initiation and propagation of GBD occurs on the steel surface when the carbon steels are polarized in the A–P region. Based on the results shown in Fig. 1 and Table 2, the electrochemical characteristic of the A–P transition, including A–P–T and A–P–P, was also observed in the PPCs for the following cases: from pH 1 to pH 4 in HNO3–NaNO2 solutions, at pH 3 and pH 4 in HCl–NaNO2 solutions and at pH 3 and pH 4 in HAc–NaNO2 solutions.

To confirm the occurrence of GBD, four steels were polarized to different potential values according to their electrochemical behaviors: in the cases of A–P–T and A–P–P, the steels were polarized to the initial passivation potential (marked in Fig. 1a and e); in the case of sP–T, the steels were polarized to the transpassivation potential (marked in Fig. 1b); in the case of sP–P, the steels were polarized to the pitting potential (marked in Fig. 1f); and in the case of A, the steels were polarized to the anodic potential value equal to the pitting potential. After this, SEM was applied to observe the surface morphologies. To facilitate comparison, the detailed results of the SEM observation are also listed in Table 2. Based on the results shown in Table 2, under the electrochemical characteristic of the A–P transition, an obvious corrosion dissolution along the GBs has been observed on the steel surface, indicating that the

| Steel | C   | Mn  | P   | S   | Si  | Cr  | Mo  | Ni  | Cu  | Al  | Fe  |
|-------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| L80   | 0.190 | 1.370 | 0.010 | 0.004 | 0.230 | 0.024 | 0.034 | 0.023 | 0.023 | 0.036 | Balance |
| N80   | 0.240 | 1.280 | 0.015 | 0.015 | 0.310 | 0.015 | 0.026 | 0.026 | 0.015 | 0.020 | Balance |
| X65   | 0.030 | 1.510 | 0.024 | 0.005 | 0.170 | 0.038 | 0.016 | 0.025 | 0.040 | 0.020 | Balance |
| Q235  | 0.160 | 0.530 | 0.015 | 0.045 | 0.300 | —    | —    | —    | —    | —    | Balance |
Occurrence of GBD is a general corrosion behavior for the L80, N80, X65 and Q235 steels, rather than a special corrosion behavior as in the case of the Q235 steel. Moreover, the presence of non-uniform passive films, mechanical scratches and corrosion pits was observed on the steel surface under the electrochemical behaviors of sP-T, A and sP-P, respectively.

Fig. 1 Potentiodynamic polarization curves of the L80, N80, X65 and Q235 steels in HNO₃–NaNO₂, HCl–NaNO₂ and HAc–NaNO₂ solutions with 0.01 M NaNO₂ and different pH values: (a) L80–HNO₃–NaNO₂, (b) N80–HNO₃–NaNO₂, (c) X65–HNO₃–NaNO₂, (d) Q235–HNO₃–NaNO₂, (e) L80–HCl–NaNO₂, (f) N80–HCl–NaNO₂, (g) X65–HCl–NaNO₂, (h) Q235–HCl–NaNO₂, (i) L80–HAc–NaNO₂, (j) N80–HAc–NaNO₂, (k) X65–HAc–NaNO₂ and (l) Q235–HAc–NaNO₂.
Table 2  Electrochemical behaviors and SEM morphologies of the L80, N80, X65 and Q235 steels polarized to different potential values in HNO$_3$–NaNO$_2$, HCl–NaNO$_2$ and HAc–NaNO$_2$ solutions with 0.01 M NaNO$_2$ and different pH values

| Acid          | pH | Steel | L80 | N80 | X65 | Q235 |
|---------------|----|-------|-----|-----|-----|------|
| HNO$_3$ + NaNO$_2$ | pH 1 | A-P-T | A-P-T | A-P-T | A-P-T | A-P-T |
|               | pH 2 | A-P-T | A-P-T | A-P-T | A-P-T | A-P-T |
|               | pH 3 | A-P-T | A-P-T | A-P-T | A-P-T | A-P-T |
|               | pH 4 | sP-T  | sP-T  | sP-T  | sP-T  | sP-T  |
|               | pH 5 | A     | A     | A     | A     | A     |
| HCl + NaNO$_2$ | pH 2 | A-P-P | A-P-P | A-P-P | A-P-P | A-P-P |
|               | pH 3 | A-P-T | A-P-T | A-P-T | A-P-T | A-P-T |
As observed from the abovementioned results, the relationship between the electrochemical characteristic of the A–P transition and the occurrence of GBD for the L80, N80, X65 and Q235 steels in HNO₃–NaNO₂, HCl–NaNO₂ and HAc–NaNO₂ solutions was very close. The appearance of the A–P transition in the PPC indicated the occurrence of GBD on the steel surface and vice versa. However, note that the effects of the acid type, pH value and steel type on the electrochemical behaviors were very significant and have been discussed hereinafter.

For the L80 steel in the pH 2 solutions, the electrochemical behaviors of A–P–T and A were respectively shown in the PPCs in the pH 4 HNO₃–NaNO₂ solution and pH 4 HCl–NaNO₂ solution, which were mainly attributed to the difference in the oxidability and the components HNO₃ and HCl. HNO₃ showed both strong oxidability and strong acidity. In contrast, HCl did not show oxidability and strong acidity. Moreover, the inhibition of NO₃⁻ and the aggression of Cl⁻ in corrosive environments were reported. NO₃⁻ promoted passivation on the steel surface; however, Cl⁻ attacked the surface passive film. Therefore, the presence of the A–P transition in the pH 4 HNO₃–NaNO₂ solution and the corresponding absence in the pH 2 HCl–NaNO₂ solution was observed. Note that the electrochemical behaviors of the four steels in the pH 1 HCl–NaNO₂ solution were similar to those in the pH 2 HCl–NaNO₂ solution. Therefore, the PPC and SEM results in the former solution are not provided herein. Because of the relatively weak acidity of HAc, the pH 1 and pH 2 HAc–NaNO₂ solutions could not be obtained. In the pH 3 and pH 4 solutions, the L80 steel showed the A–P–T, A–P–P and A–P–T behaviors in the HNO₃–NaNO₂, HCl–NaNO₂ and HAc–NaNO₂ solutions, respectively. The electrochemical characteristic of the A–P transition was due to the effectiveness of NO₂⁻ and the decrease in the H⁺

| Acid    | pH | Steel | L80 | N80 | X65 | Q235 |
|---------|----|-------|-----|-----|-----|------|
| HAc + NaNO₂ | 4  | A–P–P | A–P–P | A–P–P | A–P–P | A–P–P |
|         |    | sP–P  | sP–P  | sP–P  | sP–P  | sP–P  |
|         | 5  | A–P–T | A–P–T | A–P–T | A–P–T | A–P–T |
| HAc + NaNO₂ | 4  | A–P–T | A–P–T | A–P–T | A–P–T | A–P–T |
|         | 5  | sP–T  | sP–T  | sP–T  | sP–T  | sP–T  |
concentration; moreover, the presence of Cl\(^-\) derived from HCl was responsible for pitting.\(^8\) In the pH 5 solutions, the lack of the H\(^+\) concentration contributed to the sP-T behavior in the HNO\(_3\)–NaNO\(_2\) and HAc–NaNO\(_2\) solutions and the sP–P behavior in the HCl–NaNO\(_2\) solution for the L80 steel, which was attributed to the following reaction.\(^5\)–\(^7\)

\[
2\text{Fe}^{2+} + 2\text{OH}^- + 2\text{NO}_2^- \rightarrow 2\text{NO} + \gamma\text{-Fe}_2\text{O}_3 + \text{H}_2\text{O} \quad (6)
\]

The effect of the acid type on the electrochemical behavior for the N80, X65 and Q235 steels was similar to that in the case of the L80 steel.

In the HNO\(_3\)–NaNO\(_2\) solutions, with an increase in the pH value, the L80 steel showed the electrochemical behaviors of A–P–T in the pH 1–pH 4 solutions and sP–T in the pH 5 solution. The relatively wide pH range from pH 1 to pH 4 for the electrochemical characteristic of the A–P transition was attributed to the strong oxidizability of HNO\(_3\) and the inhibition of NO\(_3^-\). In contrast, in the HCl–NaNO\(_2\) and HAc–NaNO\(_2\) solutions, the electrochemical characteristic of the A–P transition for the L80 steel was present only in the pH 3 and pH 4 solutions due to the component HCl and the weak acidity of HAc, which has been explained in the previous discussion. In the pH 2 HCl–NaNO\(_2\) solution, the absence of the A–P transition for the L80 steel was attributed to the excess H\(^+\) concentration\(^31\) and the aggression of Cl\(^-\).\(^3\) Similarly, the effect of the pH value on the electrochemical behavior for the N80, X65 and Q235 steels was similar to that for the L80 steel.

Regarding the effect of the steel type on the electrochemical behavior, note that with the same acid type and pH value, the four steels showed the same electrochemical behavior; this indicated that the composition and alloyed element did not affect the electrochemical characteristic of the A–P transition and the occurrence of GBD. However, the different SEM morphologies were due to the different microstructures of the four steels, which have been studied further.

To further confirm the relationship between the A–P transition and GBD, the four steels were polarized to the initial passivation potential (marked in Fig. 2a) in the CO\(_2\)–NaNO\(_2\) solution and then observed by SEM. The results of the PPC test and SEM observation are shown in Fig. 2. All four steels showed the electrochemical behavior of A–P–T, and an obvious GBD was observed on the steel surface. For the L80, N80, X65 and Q235 steels, the results obtained in the CO\(_2\)–NaNO\(_2\) solution were consistent with those obtained in the HNO\(_3\)–NaNO\(_2\), HCl–NaNO\(_2\), HAc–NaNO\(_2\) and CO\(_2\)–NaNO\(_2\) solutions; GBD could be observed on the steel surface at pH 3.7, which was in the range from pH 3 to pH 4.

However, it should be clarified that in a previous study, it has been reported that the occurrence of GBD is attributed to the combined effects of CO\(_2\) and NO\(_2^-\). NO\(_2^-\) promoted passivation on the surface of the crystal grains, whereas CO\(_2\) induced dissolution in the vicinity of the grain boundaries.\(^37\) In this study, it was concluded that H\(^+\), rather than CO\(_2\), resulted in the occurrence of GBD in acidic solutions containing NO\(_2^-\). For the L80, N80, X65 and Q235 steels in the HNO\(_3\)–NaNO\(_2\), HCl–NaNO\(_2\), HAc–NaNO\(_2\) and CO\(_2\)–NaNO\(_2\) solutions, the relationship between the A–P transition and GBD has been discussed hereinafter.

GBD was observed on the steel surface when the four steels showed the electrochemical behaviors of A–P–T and A–P–P. Before the applied potential reached the transpassivation potential and the pitting potential, the PPCs of A–P–T and A–P–P were similar and composed of an activation (A) region, activation-passivation (A–P) region and passivation (P) region. Furthermore, according to previous studies\(^27\)–\(^30\) and the current results obtained in this study, the initiation and propagation of GBD occurred before the four steels were polarized in the P region. Therefore, this study was mainly focused on the anodic and cathodic reactions occurring in the A region and A–P region. Fig. 3 shows a schematic to describe the electrode reactions in the A region and the A–P region. For the L80, N80, X65 and Q235 steels in the HNO\(_3\)–NaNO\(_2\), HCl–NaNO\(_2\), HAc–NaNO\(_2\) and CO\(_2\)–NaNO\(_2\) solutions, before the applied potential reached the A–P transition potential, the dominant anodic and cathodic reactions were Fe oxidation (eqn (5)) and H\(^+\) reduction (eqn (1)), respectively. With a positive shift of the applied potential, the electrode reactions accelerated. Therefore, the anodic current density gradually increased with potential in the A region until the applied potential reached the A–P transition potential. However, due to the presence of NO\(_2^-\) and the continuously positive shift of the applied potential, the following cathodic reactions involving the NO\(_2^-\)/HNO\(_2\) reduction occurred:

\[
2\text{NO}_2^- + 8\text{H}^+ + 6\text{e}^- \rightarrow \text{N}_2 + 4\text{H}_2\text{O} \quad (7)
\]

\[
2\text{HNO}_2 \rightarrow \text{N}_2\text{O}_4 + 2\text{H}^+ + 2\text{e}^- \quad (8)
\]

\[
2\text{NO}_2^- + 6\text{H}^+ + 4\text{e}^- \rightarrow \text{N}_2\text{O}_3^- + 3\text{H}_2\text{O} \quad (9)
\]

\[
\text{HNO}_2 + \text{H}_2\text{O} \rightarrow \text{NO}_3^- + 3\text{H}^+ + 2\text{e}^- \quad (10)
\]

Moreover, the anodic reaction of Fe\(^{2+}\) oxidation may be possible:

\[
\text{Fe}^{2+} \rightarrow \text{Fe}^{3+} + \text{e}^- \quad (11)
\]

The reason is that the \(E_{\text{st}}\) of NO\(_2^-\)/HNO\(_2\) reductions (eqn (7)–(10)) is significantly positive than that of the Fe\(^{2+}\) oxidation (eqn (11)). However, kinetic studies and potential calculations are necessary to verify the actual cathodic reactions of the NO\(_2^-\)/HNO\(_2\) reduction. It is generally accepted that the electrochemical characteristic of the A–P transition is very closely associated with the generation of Fe\(^{2+}\), which is the main component in the surface passive film.\(^39\)–\(^44\) With a positive shift of the applied potential, when the applied potential reached up to the A–P transition potential, the anodic current density gradually decreased with potential in the A–P region; this suggested the activation of Fe\(^{2+}\) oxidation and the formation of the passive film. Furthermore, the cathodic reactions for the NO\(_2^-\)/HNO\(_2\) reduction played a critical role in the abovementioned process.\(^28\)–\(^31\) On the other hand, the metallurgical phenomenon of the grain boundary segregation (GBS) was generally present.
on the alloy steels and carbon steels, which resulted in the difference between passivation capabilities of GIs and GBs. In a previous study, it has been reported that GIs containing high level of Fe showed better passivation capability than GBs with high levels of Si and Mn. Therefore, when the applied potential moved to the A–P region, the electrochemical micro-couple between the GIs and GBs was established spontaneously to induce the occurrence of GBD. However, between the GIs and the GBs, the detailed mechanisms about the establishment of the electrochemical micro-couple and the driving force for the potential difference need further investigation.

Fig. 2 Potentiodynamic polarization curves and surface SEM morphologies of the L80, N80, X65 and Q235 steels in a CO$_2$–NaNO$_2$ solution: (a) potentiodynamic polarization curves, (b) L80 SEM image, (c) N80 SEM image, (d) X65 SEM image and (e) Q235 SEM image.
4. Conclusions

In this study, four steels (L80, N80, X65 and Q235) were investigated in four acidic solutions (HNO₃, HCl, HAc and CO₂) containing NO₂⁻ to reveal the relationship between the electrochemical characteristic of the A-P transition and the occurrence of GBD. The main conclusions were as follows:

(1) The relationship between the A-P transition and GBD was very close. The appearance of the A-P transition in the PPC indicated the occurrence of GBD on the steel surface. Moreover, it was confirmed that under the electrochemical characteristic of the A-P transition, the occurrence of GBD was a general corrosion behavior for carbon steels and alloy steels in acidic media containing NO₂⁻.

(2) The effect of the steel type on the electrochemical characteristic of the A-P transition and the occurrence of GBD was not obvious. However, the effects of acid type and pH value were very significant. The A-P transition and the GBD were present from pH 1 to pH 4 in the HNO₂–NaNO₂ solutions, at pH 3 and pH 4 in the HCl–NaNO₂ solutions and at pH 3 and pH 4 in the HAc–NaNO₂ solutions.

Conflicts of interest

There are no conflicts to declare.

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