Effects of pre-oxidation on the corrosion behavior of pure Ti under coexistence of solid NaCl deposit and humid oxygen at 600 °C: the diffusion of chlorine

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The effect of pre-oxidation on the corrosion behavior of pure Ti covered with a solid NaCl deposit in the humid O₂ flow at 600 °C is studied. The oxide scale, formed by pre-oxidation, protects the substrate from the NaCl induced corrosion during the initial stage. However, the corrosion of the pre-oxidized sample is severely accelerated by solid NaCl after an incubation period. The chlorine, generated from the decomposition of solid NaCl, diffuses into the oxide/substrate interface as ions during the incubation period, which was observed by ToF–SIMS. The chlorine at the oxide/substrate interface induces the fast corrosion after the incubation period although the pre-oxidation scale is complete and compact.

Compressor blades of airplanes and ships suffer severe corrosion in marine environment, which is mainly due to the fact that marine air contains abundant salts (especially NaCl) and water vapor. At the temperature of 300–600 °C where compressor blades operate, NaCl appears in solid state and gets deposited on the metal surface. The corrosion of several metals and alloys of compressor blades gets accelerated under a synergistic effect of the solid NaCl deposit and humid oxygen.

Extensive works have been done on the corrosion behavior of several metals and alloys of compressor blades. The authors in paper¹¹ demonstrated that the corrosion behavior of pure Fe, pure Cr, Fe–Cr alloy, Ti alloy and Ni alloy at 600 °C under a deposit of solid NaCl deteriorated sharply. Similarly, the authors investigated the influence of KCl on the oxidation of the 304-type (Fe18Cr10Ni) austenitic stainless steel at 600 °C in 5% O₂ and in 5% O₂ + 40% H₂O in paper¹²⁻¹⁷, and the results showed that small additions of potassium chloride strongly accelerated high temperature corrosion. The fast corrosion is mainly due to a chemical reaction of the solid NaCl/KCl deposit with oxides. This reaction destroys the protective oxide scale on the surface and forms a porous non-protective scale (i.e., the scale has numerous holes and voids¹,³,⁶,⁸,¹³,¹⁷). It is also reported in these investigations that the key factor for the fast corrosion is the chlorine, HCl or Cl₂, which cyclically reacts with the substrate at the oxide/substrate interface. The gaseous molecule of chlorine, which is formed during the chemical reaction of solid NaCl and oxide¹,³,⁶,¹₈, differs inward quickly through macro defects (like cracks and holes) in the scale of corrosion products. The chlorine then reacts cyclically with the substrate to generate volatile products (for example CrCl₃) which break the protective scale¹,³,⁶,⁸.

In our previous work¹¹, we observe that the scale of corrosion products of Ti60 alloys can be divided into an outer the corrosion layer and an inner corrosion layer after exposure at 600 °C in the environment of NaCl + H₂O + O₂. The chlorine exists in a Ti-Cl bond which is formed by replacing O with Cl within lattices of

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Ti oxides of the inner corrosion layer. We think the chlorine can diffuse inward as ion. In this case, Cl\(^-\) comes from the decomposition of solid NaCl and the residual sodium reacts with oxides on the surface to form metallic acid salts (like Na\(_2\)CrO\(_4\) and Na\(_4\)Ti\(_5\)O\(_{12}\)). In order to make clear the corrosion mechanism, we need to first figure out the state of the chlorine and study its diffusion in the scale of corrosion products, especially during the destruction process of the scale of protective oxide, like TiO\(_2\) and Cr\(_2\)O\(_3\).

When pure Ti is oxidized in the pure O\(_2\) condition at 600 °C, a compact and even oxide scale forms on the surface. This oxide scale can simulate the passive film to protect the substance from the corrosive environment. When the pre-oxidized pure Ti is exposed in the NaCl + H\(_2\)O + O\(_2\) environment, this oxide scale can slow down the destruction process of the protective passive film such that the behavior of the solid NaCl can be investigated in detail. Thus, to clarify the micro-mechanism under the specific environment of NaCl + H\(_2\)O + O\(_2\), we need to study the effects of pre-oxidation on the corrosion behavior of pure Ti.

In this paper, we investigate the effects of pre-oxidation on the corrosion behavior of pure Ti underneath a solid NaCl deposit in humid O\(_2\) flow at 600 °C, using scanning electron microscope (SEM) equipped with an energy dispersive spectrometer (EDS), X-ray diffraction (XRD) and time of flight-secondary ion mass spectrometry (ToF–SIMS). After examining the detailed properties of the corrosion products and the diffusion of chlorine in the oxide scale, we discuss possible micro acceleration mechanisms of the chlorine based on experiment results.

Result

Corrosion kinetics. The mass gain curves of pure Ti under different conditions are shown in Fig. 1. For the pre-oxidized samples, the mass gain under the condition of H\(_2\)O + O\(_2\) (about 0.17 mg/cm\(^2\)) is slightly larger than that under the condition of O\(_2\) (about 0.09 mg/cm\(^2\)). In fact, under both the above two conditions the corrosion is minor. In the initial stage (incubation period), the mass gain is slight when the pre-oxidized pure Ti samples under NaCl deposit. In contrast, the mass gain increases rapidly after this incubation period. After 20 h exposure in NaCl + H\(_2\)O + O\(_2\) or NaCl + O\(_2\), the mass gains are about 1.8 mg/cm\(^2\) and 1.4 mg/cm\(^2\), respectively, which are about one order of magnitude larger than that in the absence of the solid NaCl deposit. Thus, we reach the conclusion that the solid NaCl accelerates the corrosion of the pre-oxidized pure Ti in O\(_2\) or humid O\(_2\).

For the bare samples, the mass gain increases dramatically during the whole exposure in NaCl + H\(_2\)O + O\(_2\). After exposed for 20 h, the mass gain is about 6.0 mg/cm\(^2\), which is increased by 3.5 times compared with the pre-oxidized samples. Therefore, we can conclude that the pre-oxidation improves the corrosion resistance of pure Ti coated with a solid NaCl deposit in humid O\(_2\) at 600 °C, especially in the incubation period.

Phase composition and microstructure of the pre-oxidized sample. For the pure Ti samples after pre-oxidation in O\(_2\) flow for 20 h, the surface and the cross-sectional morphologies are shown in Fig. 2. As can be observed in Fig. 2a,b, a continuous, compact and even oxide scale is formed on the surface. The thickness of the oxide scale is about 1 μm (Fig. 2b). We show the results of EDS analysis on the oxide scale in Fig. 2c. As can be seen, it consists of Ti and O. Through XRD analysis (which is not shown here), the phase composition of the pre-oxidized sample surface is only TiO\(_2\).

Phase composition and microstructure of the hot corrosion products forming on the bare samples. For the bare samples after 20 h exposure in NaCl + H\(_2\)O + O\(_2\), the surface and the cross-sectional morphologies are shown in Fig. 3. As can be observed, the scale of corrosion products contains two parts: one with a compact structure, and the other with a grid structure part. The corrosion products with a compact structure (marked in Fig. 3a,b) are very complete, uniform and compact. The corrosion products with a grid structure
(marked in Fig. 3a,b) contain lots of holes (filled with Ni plating as shown in Fig. 3b), which penetrate the corrosion scale. These through-holes are the fast diffusion path for the corrosion medium. Below the corrosion scale, a loose layer, containing some holes (close pores) and cracks, is formed within the substrate due to the outward diffusion of Ti. The thickness of the corrosion product scale and the loose layer is about 45 μm and 15 μm, respectively.

In Fig. 4, we present the XRD patterns of the bare samples exposed in NaCl + H₂O + O₂ up to 20 h. The results show that the corrosion products mainly consist of TiO₂ and some Na₄Ti₅O₁₂, with the residual NaCl.

**Phase composition and microstructure of the hot corrosion products forming on the pre-oxidized samples.** For the pre-oxidized pure Ti samples after 100 h exposure in dry O₂ flow, the surface and the cross-sectional morphologies are shown in Fig. 5. As can be seen in Fig. 5a,b, a very thin, compact and continuous scale is formed on the pre-oxidized samples after 100 h exposure in dry O₂ flow. From Fig. 5c, we can see that the thickness of the oxide scale is about 2.5 μm.

For the pre-oxidized samples after 100 h exposure in humid O₂ flow, the surface and the cross-sectional morphologies are presented in Fig. 6. As can be observed from Fig. 6a,b, a very thin, compact and continuous scale which contains both granulated corrosion products and compact products was formed on the pre-oxidized
Figure 4. X-ray diffraction patterns of the bare samples exposed in NaCl + H₂O + O₂ up to 20 h at 600 °C.

Figure 5. Surface morphologies (a), the high magnification image (b) and cross-sectional morphologies (c) of the pre-oxidized samples after oxidized in dry O₂ at 600 °C for 100 h.
samples. From Fig. 6c, we can see that the oxide scale is divided into a loose outer layer and a compact inner layer, with a thickness of about 1.2 μm and 1 μm, respectively.

For the pre-oxidized samples after 20 h exposure in NaCl + O₂, the surface and the cross-sectional morphologies are displayed in Fig. 7. As can be seen in Fig. 7a, the corrosion products on the surface are in clusters. As can be observed in Fig. 7b,c, the corrosion product scale can be divided into an outer layer and an inner layer. The outer layer is filled with clustered corrosion products, and the inner layer whose thickness is about 5 μm contains loose corrosion products.

For the pre-oxidized samples after 20 h exposure in NaCl + H₂O + O₂, the surface and the cross-sectional morphologies are depicted in Fig. 8. As can be observed in Fig. 8a,b, corrosion products have a very thin, compact and continuous scale, with some blind holes forming on the surface where we can also observe some defects (like pore). Furthermore, the scale can be divided into a loose inner corrosion layer and a compact outer corrosion layer. As shown in Fig. 8c, the thickness of the outer corrosion layer and the inner corrosion layer is 8 μm and 10 μm, respectively.

For the pre-oxidized samples after 1 h exposure in NaCl + H₂O + O₂, the surface and the cross-sectional morphologies are shown in Fig. 9. From Fig. 9a, we can observe a compact scale on which some loose corrosion products are scattered. In Fig. 9b, we can observe that under the loose product, the scale still remains complete and compact. This implies that the loose products are formed due to the reaction of the outward diffusion metal ions with the corrosive species. From Fig. 9d, we can observe that the thickness of this scale is about 1 μm.

For the pre-oxidized samples after 5 h exposure in NaCl + H₂O + O₂, the surface and the cross-sectional morphologies are depicted in Fig. 10. From Fig. 10a, we can observe that the porous corrosion products spread over the entire surface. This is consistent with the fact in Fig. 1 that the mass gain increased rapidly after the
The incubation period and the corrosion is severe. From Fig. 10b, we can observe that the scale of the corrosion products is compact and its thickness is about 1.2 μm. We can also observe some loose parts at the interface between the scale and metal.

The corrosion products of the pure Ti samples are identified by XRD. The XRD patterns in Fig. 11 show that only TiO₂ was formed on the pre-oxidized samples after 100 h exposure in dry O₂ flow (Fig. 11a) or in H₂O + O₂ flow (Fig. 11b). In Fig. 12, we can observe that the corrosion products are TiO₂, Ti₂O and Na₄Ti₅O₁₂ on the pre-oxidized samples after 20 h exposure in NaCl + O₂ (Fig. 12b), whereas the Na₄Ti₅O₁₂ cannot be identified by XRD after 5 h exposure (Fig. 12a). As shown in Fig. 13c, after the pre-oxidized samples are exposed in NaCl + H₂O + O₂ for 20 h, the corrosion products contain both TiO₂ and Na₄Ti₅O₁₂. There still remains residual NaCl as well as the substrate (pure Ti) on the surface. As shown in Fig. 13b, after 5 h exposure in NaCl + H₂O + O₂ (which is after the incubation period), the corrosion products also contain Na₄Ti₅O₁₂. Since the diffracted intensity of Na₄Ti₅O₁₂ in Fig. 13c are larger than that in Fig. 13b, we can infer that more Na₄Ti₅O₁₂ is formed when the pre-oxidized samples are exposed in NaCl + H₂O + O₂ for a longer time period. As shown in Fig. 13a, after 1 h exposure in NaCl + H₂O + O₂ (which is during the incubation period), there are not Na₄Ti₅O₁₂ among the corrosion products. We conclude the corrosion products under the above different conditions in Table 1.

Figure 7. Surface morphologies (a) and cross-sectional morphologies (b) and the high magnification image (c) of the pre-oxidized samples after corrosion in NaCl + O₂ at 600 °C for 20 h.

Figure 8. Surface morphologies (a) and cross-sectional morphologies (b) and the high magnification image (c) of the pre-oxidized samples after corrosion in NaCl + H₂O + O₂ at 600 °C for 20 h.
Figure 14 shows the depth profiles for the negative ions from the pre-oxidized samples after 1 h and 5 h exposure in NaCl + H2O + O2. The O$_2^-$ ion signal in Fig. 14 is applied here to indicate the distribution of oxygen in the scale and to identify the interface between the corrosion products scale and the substrate. In Fig. 14a, the O$_2^-$ ion exhibits two levels of concentration, and the transition region is around the interface of the corrosion product scale and the substrate. Therefore, the average of logarithmic value is pinpointed as the interface as shown in Fig. 14a. Similarly, this interface is also identified in Fig. 14b.

In Fig. 14a, we can also observe that after 1 h exposure in NaCl + H2O + O2, Cl$^-$ ions first enrich on the surface of the pre-oxidized samples (which was caused by the residual NaCl) and then decrease rapidly in the corrosion product scale. Afterwards, Cl$^-$ is enriched at the oxide/metal interface, which indicates that the chlorine penetrates the oxide and accumulates at the oxide/metal interface. In Fig. 14b, after the pre-oxidized samples are exposed in NaCl + H2O + O2 for 5 h, a similar distribution of Cl$^-$ in the corrosion product scale was observed. Particularly, the Cl$^-$ ion is still enriched when the O$_2^-$ ion decreases around the oxide/metal interface, which implies that some chlorine diffuses into the metal substrate. The intensity of Cl$^-$ around the oxide/metal interface is stronger in Fig. 14b than that in Fig. 14a. This implies that more chlorine penetrates into the oxide after a longer exposure in the presence of solid NaCl.

The OH$^-$ signal in Fig. 14 is applied here to indicate the distribution of H in the scale. From Fig. 14a, the distribution of the OH$^-$ is similar with Cl$^-$. First, the OH$^-$ ion enriches on the surface and then decrease rapidly in the corrosion product scale. This is the information of water vapor. Afterwards, OH$^-$ is enriched at the oxide/metal interface where Cl$^-$ is accumulated, which indicates that the H penetrates the oxide along with the chlorine.
Figure 10. Surface morphologies (a) and cross-sectional morphologies (b) of the pre-oxidized samples under a solid NaCl deposit layer in humid O₂ for 5 h at 600 °C.

Figure 11. X-ray diffraction patterns of the pre-oxidized samples after exposed in dry O₂ flow (a) and in humid O₂ flow (b) for 100 h.

Figure 12. X-ray diffraction patterns of the pre-oxidized samples after exposed in NaCl + O₂ for 5 h (a) and for 20 h (b).
The acceleration of NaCl to the corrosion of pure Ti.

As shown in Fig. 2, after pure Ti samples were oxidized in the pure O₂ flow at 600 °C for 20 h, a compact TiO₂ scale with a thickness of about 1 μm was formed on the surface, which is caused by the following reaction between the metal substrate and O₂:

\[ \text{Ti} + \text{O}_2 = \text{TiO}_2 \]  

In contrary, the mass gain of pure Ti under a solid NaCl deposit layer (NaCl + H₂O + O₂) increased quickly and the mass gain is about 6.0 mg/cm² after 20 h exposure (Fig. 1), which is much larger than that in the absence of solid NaCl (about 0.3 mg/cm²). The resulting scale consists of plentiful and porous corrosion products (Fig. 3) and its thickness is about 45 μm. The above phenomena indicate that pure Ti suffered severe corrosion when exposed in NaCl + H₂O + O₂. This severe corrosion has been discussed in our previous work and the main reason is that the protective TiO₂ scale cannot be formed due to the occurrence of a series of chemical reactions, and the corrosion becomes active. In the previous mechanism, NaCl first destroys the protective scale as follow:

\[ \text{NaCl} + \text{TiO}_2 = \text{Na}_4\text{Ti}_5\text{O}_{12} + \text{4HCl} \]  

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Figure 13. X-ray diffraction patterns of the pre-oxidized samples after exposed in NaCl + H₂O + O₂ for 1 h (a), 5 h (b) and 20 h (c).

Table 1. Corrosion products forming on the bare samples and the pre-oxidized samples after hot corrosion.

| Samples          | Environment          | Corrosion time | 1 h | 5 h | 20 h | 100 h |
|------------------|----------------------|----------------|-----|-----|------|-------|
| Bare samples     | NaCl + H₂O + O₂      |                | –   | –   | TiO₂ | –     |
|                  | O₂                   |                | –   | –   | –    | –     |
| Pre-oxidized     | NaCl + H₂O + O₂      | TiO₂, Ti₂O    | TiO₂, Ti₂O, Na₄Ti₅O₁₂ | TiO₂, Ti₂O, Na₄Ti₅O₁₂ | –   |
|                  | NaCl + O₂            | –             | TiO₂, Ti₂O, Na₄Ti₅O₁₂ | TiO₂, Ti₂O, Na₄Ti₅O₁₂ | –   |
|                  | H₂O + O₂             | –             | –   | –   | TiO₂ | –     |
|                  | O₂                   | –             | –   | –   | –    | –     |
The TiCl₄ reacts with H₂O to form HCl:

\[ \text{TiCl}_4 + 2\text{H}_2\text{O} \rightarrow \text{TiO}_2 + 4\text{HCl} \]  

Then, HCl reacts with the substrate cyclically:

\[ 2\text{HCl} + \text{Ti} = \text{TiCl}_2 + \text{H}_2 \]  
\[ 4\text{HCl} + \text{Ti} = \text{TiCl}_4 + 2\text{H}_2 \]  
\[ 2\text{TiCl}_2 + \text{O}_2 + 2\text{H}_2\text{O} = 2\text{TiO}_2 + 4\text{HCl} \]  
\[ 2\text{H}_2 + \text{O}_2 = 2\text{H}_2\text{O} \]

These reactions are thermodynamically spontaneous due to the negative ΔG* at 600 °C, as presented in Table 2. As shown in Fig. 3b, in the corrosion product scale there are holes which provide rapid diffusion channels for corrosive species (for example, oxygen, chlorine and water vapor). Meanwhile, a corrosion scale is formed by the outward diffusion of Ti which results in a 15 μm loose layer that is observed under the corrosion products. Hence, the corrosion product scale is non-protective and the corrosion of pure Ti is greatly accelerated by solid NaCl in an O₂ + H₂O environment at 600 °C, which is similar with the observation for several Ti alloy11,18–20.

**The protection of pre-oxidation scale to the pure Ti.** After pre-oxidation in O₂ flow for 20 h, a continuous, compact and even oxide scale is formed on the surface (Fig. 2). The pre-oxidation scale is only TiO₂ (Fig. 2c) and its thickness is about 1 μm (Fig. 2b). Since this scale can protect the substrate, the mass gain of the pre-oxidized samples is small in the dry O₂ or humid O₂ (Fig. 1).

When pre-oxidized samples are exposed in the presence of solid NaCl (NaCl + O₂ and NaCl + H₂O + O₂), the mass gain of the pre-oxidized samples is smaller than that of the bare samples (Fig. 1), especially in the incubation period (the mass gain increases slowly). The resulting scale on the pre-oxidized samples (which is compact as

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**Table 2.** Standard Gibbs free energy changes of reactions at 600 °C. * The G* value of Na₄Ti₅O₁₂ was calculated by the first principles. And others were calculated by the HSC Chemistry.

| Reaction | ΔG* (kJ/mol) |
|----------|--------------|
| Ti + O₂ → TiO₂ | −785 |
| 4NaCl + 6TiO₂ → Na₄Ti₅O₁₂ + TiCl₄ | −65 |
| 4NaCl + 5TiO₂ + 2H₂O → Na₄Ti₅O₁₂ + 4HCl | −195 |
| TiCl₂ + 2H₂O → TiO₂ + 4HCl | −130 |
| Ti + 2HCl → TiCl₂ + H₂ | −174 |
| Ti + 4HCl → 2TiCl₂ + 4HCl | −256 |
| 2TiCl₂ + 2H₂O + O₂ → 2TiO₂ + 4HCl | −905 |
| 2H₂ + O₂ → 2H₂O | −399 |

**Figure 14.** ToF–SIMS sputter depth profiles in the corrosion products scales formed on the pre-oxidized samples under a NaCl deposit in an atmosphere of humid O₂ at 600 °C for 1 h (a) and 5 (b).
shown in Figs. 7 and 8) is thinner than that on the bare samples (which is porous as shown in Fig. 3). Thus, the pre-oxidation can protect the pure Ti samples from the corrosion under NaCl deposit. This is mainly attributed to the compact TiO₂ scale formed on the surface in the pre-oxidation (Fig. 2). During the pre-oxidation, the thickness of the formed TiO₂ scale is about 1 μm, and the pre-oxidation scale is larger thicker than the passive film forming on the bare sample. Thus, the TiO₂ scale can be a barrier layer for the diffusion of corrosive species (for example, oxygen, water vapor and HCl) and reduce the corrosion to a certain extent, especially in the incubation period.

The destruction of pre-oxidation scale. When exposed in humid O₂, the pre-oxidized samples suffer a slightly more serious oxidation than when exposed in dry O₂ (Fig. 1). This is because H₂O tends to dissociate at the defects on TiO₂/(110) planes into free H atoms and OH⁻ groups when the pre-oxidized samples is exposed in humid O₂. The papers reported that along the crystal channels in c-axis direction, the diffusion rate of hydrogen is at least one order of magnitude larger than that in the perpendicular direction of c-axis. Thus, the generated hydrogen atoms are more prone to diffuse into TiO₂ through the former one. This dissolution of hydrogen in TiO₂ will form hydrogen defects, thus the hydrogen can increase the concentration of crystal defects and increase the outward diffusion of Ti ions. This fast diffusion in the scale causes a larger corrosion rate and hence the hole forms in the oxide scale (Fig. 6), which make the corrosion product scale thicker and more porous (Fig. 6c). Thus, the water vapor slightly reduces the protection of the pre-oxidation scale by forming the H defects.

From Fig. 1, we can observe the following: (1) after the incubation period, the mass gain of the pre-oxidized samples increases more rapidly when exposed in the presence of the solid NaCl (NaCl + O₂ or NaCl + H₂O + O₂) than when exposed in the absence of the solid NaCl (O₂ or H₂O + O₂); (2) after 20 h exposure, the pre-oxidized samples has a larger mass gain under the condition of NaCl + O₂ or NaCl + H₂O + O₂ than under the condition of O₂ or H₂O + O₂. Additionally, we can observe that the corrosion product scale of the pre-oxidized samples is thicker when exposed in the presence of the solid NaCl (Figs. 7 and 8) than when exposed in the absence of the solid NaCl (Figs. 5 and 6). Thus, we can claim that the formation of the pre-oxidized samples is accelerated by solid NaCl. The fast corrosion and the formation of Na₄Ti₅O₁₂ on the surface (Fig. 9c,d) indicate that the “protective oxide” formed in the pre-oxidation is not completely inert to solid NaCl.

The diffusion of the chlorine in the pre-oxidation scale. The evidence from XRD and SEM/EDX show that Na₄Ti₅O₁₂ formed on the surface (see Figs. 4 and 9c,d). This indicated that the solid NaCl reacted with the TiO₂ scale. Similar observations have been reported for Ti alloy using the same experimental conditions. In that case, it was argued that the fast corrosion in an O₂ + H₂O environment is triggered by titanate formation. Thus, in an O₂ + H₂O + NaCl environment, the formation of titanate destroys the protective scale formed on the surface, and then the gathered chlorine circularly reacts with the substrate. Thus, for the corrosion of pure Ti/Ti alloy in the NaCl + H₂O + O₂ environment, the chloride is the main factor.

For the pre-oxidized samples, a thick TiO₂ scale is formed on the surface; thus, it takes more time to destroy the protective TiO₂ scale by Na₄Ti₅O₁₂ formation and the mass gain changes little during the incubation period (Fig. 1). Furthermore, after the sodium forms Na₄Ti₅O₁₂, the gathered chloride diffuses inward to the metal/oxide interface (Fig. 14a). Shu et al. report that the chloride permeates into the corrosion product scale as molecule, like HCl or Cl₂. For the pre-oxidized samples, a compact TiO₂ formed on the surface protects the substrate from the inward diffusion of chlorine and the fast corrosion. Nevertheless, during the incubation period, the chloride also diffuses into the metal/oxide interface (Fig. 14a) while the TiO₂ scale is complete (Fig. 9). This implies that the chloride cannot be molecule. In our previous work, the chloride forms Ti-Cl bond in the inner corrosion layer by the dopant Cl replacing O in the Ti oxides lattice when Ti60 alloy was exposed in the NaCl + H₂O + O₂. On the other hand, the diffusion of ion in the TiO₂ is fast and some NaCl is split into ions (Na⁺ and Cl⁻) at 600 °C. Hence, we think the chloride permeates into the corrosion product scale as ion (Cl⁻).

As shown in Fig. 14b, when the pre-oxidized samples suffer from severe corrosion in NaCl + H₂O + O₂, the chloride is also observed around the metal/oxide interface by SIMS. Additionally, the intensity of chlorine is still large when the intensity of oxygen decreases to a relatively small value. As aforementioned, the substrate locates at the places where the intensity of oxygen is small. Hence, we can infer that the chlorine diffuses into the substrate and directly binds with the metal of the substrate during the fast corrosion. The gathered compound diffuses outward, resulting in the loss of metal in the substrate. After 20 h exposure, a mass of the metal reacts with the chloride and diffuses outward, and the inner corrosion layer is formed under the outer corrosion layer (Fig. 8b).

Conclusion
During the pre-oxidation, a compact and thick TiO₂ scale forms on the surface of pure Ti. This TiO₂ scale is the barrier layer for the diffusion of corrosive species (for example, oxygen, chlorine, and water vapor), which can protect the substrate from fast corrosion. However, the “protective oxide” is not completely inert to solid NaCl and the corrosion rate of the pre-oxidized sample is also greatly accelerated by solid NaCl after an incubation period. During the incubation period, the chlorine, generated in the decomposition of solid NaCl at 600 °C, diffuses into the oxide/substrate interface as ions to enhance the corrosion rate, and then the residual sodium reacts with the oxide on the surface. After the incubation period, the chlorine diffuses into the substrate and directly reacts with the substrate, resulting in the fast corrosion.
was about 3.2 cm, the flow rate of pure O₂ was about 140 mL/min in this study) to produce the test atmosphere (humid O₂). According to the relationship between the vapor pressure of water and its temperature, we precisely first condition was with a solid NaCl deposit layer in a humid O₂ flow at 600 ℃ (denoted as NaCl + H₂O + O₂); of the furnace tube.

In this study, the temperature of the distilled water was about 70 ℃, producing about 30.8 vol.% water vapor. In order to avoid the water vapor condensing inside the thermo-balance, a counter flow of pure N₂ was passed through the thermo-balance, whose flow rate was about 400 mL/min. After the furnace reached 600 ℃ and the gas flows of humid O₂ as well as pure N₂ stabilized, the samples were quickly lowered into the constant temperature zone of the furnace tube.

In this study, the corrosion experiments were respectively carried out under the following four conditions: the first condition was with a solid NaCl deposit layer in a humid O₂ flow at 600 ℃ (denoted as NaCl + H₂O + O₂); the second condition was carried out in a humid O₂ flow at 600 ℃ (denoted as H₂O + O₂); the third condition was with a solid NaCl deposit layer in a dry O₂ flow at 600 ℃ (denoted as NaCl + O₂); and the fourth condition was carried out in a dry O₂ flow at 600 ℃ (denoted as O₂). We present the explicit environmental parameters in Table 3.

| Environment      | Mass of NaCl (mg/cm²) | Water pressure (kPa) | Flow rate of O₂ (mL/min) | Temperature (℃) |
|------------------|------------------------|----------------------|--------------------------|-----------------|
| NaCl + H₂O + O₂  | 4.0                    | 31                   | 140                      | 600             |
| NaCl + O₂        | 4.0                    | 0                    | 140                      | 600             |
| H₂O + O₂         | 0                      | 31                   | 140                      | 600             |
| O₂               | 0                      | 31                   | 140                      | 600             |

Table 3. Experimental parameters.

**Experiment**

**Materials preparation.** The material used in this study was pure Ti and the samples were cut into the size of 10 mm × 15 mm × 2.5 mm. Prior to experiments, all samples were mechanically grinded with 800# SiC paper, ultrasonically degreased in alcohol for about 20 min, and dried in the air. The bare samples were then exposed in the pure O₂ flow at 600 ℃ for 20 h for the pre-oxidization. The surfaces of preheated samples were covered with a layer of NaCl deposit by repeatedly brushing and drying with saturated NaCl solution, until about 4 ± 0.2 mg/cm² solid NaCl was deposited on.

**Corrosion experiments.** The corrosion tests were carried out in a thermo-balance. The continuous mass gain during the corrosion experiment was obtained with a thermo-gravimetric analysis (TGA).

The pure O₂ was bubbled into the distilled water with a glass bubbler (when the inner diameter of the tube was about 3.2 cm, the flow rate of pure O₂ was about 140 mL/min in this study) to produce the test atmosphere (humid O₂). According to the relationship between the vapor pressure of water and its temperature, we precisely set up the temperature of the distilled water in the glass bubbler to control the amount of the water vapor. In this study, the temperature of the distilled water was about 70 ℃, producing about 30.8 vol.% water vapor. In order to avoid the water vapor condensing inside the thermo-balance, a counter flow of pure N₂ was passed through the thermo-balance, whose flow rate was about 400 mL/min. After the furnace reached 600 ℃ and the gas flows of humid O₂ as well as pure N₂ stabilized, the samples were quickly lowered into the constant temperature zone of the furnace tube.

In this study, the corrosion experiments were respectively carried out under the following four conditions: the first condition was with a solid NaCl deposit layer in a humid O₂ flow at 600 ℃ (denoted as NaCl + H₂O + O₂); the second condition was carried out in a humid O₂ flow at 600 ℃ (denoted as H₂O + O₂); the third condition was with a solid NaCl deposit layer in a dry O₂ flow at 600 ℃ (denoted as NaCl + O₂); and the fourth condition was carried out in a dry O₂ flow at 600 ℃ (denoted as O₂). We present the explicit environmental parameters in Table 3.

**Morphologies and chemical composition analysis.** The surface morphologies and the cross-sectional morphologies of corrosion products scale were collected by SEM–EDS. The chemical composition of corrosion products was identified by XRD.

To protect the oxide scale from fracture and spall during the metallographic preparation, the corroded samples were wrapped into a thin nickel foil by electroless plating and were embedded into the epoxy resin. Then, the corroded samples were ground to 3000 grit with SiC paper. Finally, the corroded samples were polished with diamond paste. The samples were washed with distilled water to remove the residual solid NaCl, and then dried in air, before surface investigation by SEM. Under the condition of H₂O + O₂ and O₂, the mass gains of the pre-oxidized sample were very small (Fig. 1) and the corrosion was minor. The surface and the cross-sectional morphologies after a longer corrosion time (100 h) were investigated.

**Time of flight-secondary ion mass spectrometry, ToF–SIMS.** The pre-oxidized samples were first exposed in either NaCl + H₂O + O₂ or NaCl + O₂, Then, they are ultrasonically washed with distilled water to remove the residual solid NaCl. After the pre-oxidized samples were further dried in the air, they were analyzed using a ToF–SIMS5 instrument (ION-TOF GmbH), which allowed parallel mass registration with high sensitivity and high mass resolution. A cesium liquid–metal ion (LMI) gun at 20 keV beam energy was used for spatially resolved ToF–SIMS analysis.

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Author contributions L.F. designed and performed experiment. L.F., L.L., C.X.Y, A.Q.F, J.T.Y, F.H.W. and Y.L. discussed and analyzed data. L.F. and L.L. wrote the manuscript. L.L., Y.H.L., H.W revised the paper.

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