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Water vapor oxidation of ferritic 441 and austenitic 316L stainless steels at 1100 °C for short duration

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Abstract

A ferritic 441 and an austenitic 316L steels have been exposed to wet argon at 1100 °C. This study focus on the characterization of the oxide scales formed after different exposure times in the range of 2.5-20 min. Raman spectroscopy, XRD, SEM and XPS have been used. For all exposure times, 316L forms a breakaway type thick oxide scale (rupture of the pre-existing passivating film) with iron oxides on its outer part and a mix of spinels with Fe, Cr and Ni for its inner part. For lower water vapor partial pressure, iron oxides are constituted of wüstite. For higher water vapor partial pressure, iron oxides are constituted of a layer of hematite over a layer of magnetite slightly enriched in chromium. Due to strong oxidation condition, oxide scale is not always homogeneous and iron oxides spallation may occur. For 2.5 min of oxidation on 441, a very thin layer of protective chromium oxide is formed. For longer exposure time, an almost homogeneous and much thicker layer mainly consisting of Cr₂O₃ is produced. The thickness varies slightly and gradually from 4 to 20 min of oxidation. There are Mn-Cr spinels mixed with the chromium oxide. The most external part is strongly enriched in Mn and Fe in a spinel structure. The diffusivity of chromium is regarded as the main cause of the difference of oxidation behavior. In both cases, the first step is a very thin chromium oxide layer. When the oxidation conditions becomes too strong in terms of exposure time or water vapor partial pressure, this oxide layer breaks. The ferritic steel is able to heal and thicken its protective chromium oxide, preventing the breakaway. The healing would be due to the high diffusivity of chromium. The thickening would be caused by the presence of the Mn-Cr spinels which are a less effective diffusion barrier. The lower diffusivity of chromium in austenite promotes the breakaway.

Keywords: stainless steel – oxidation - water vapor - high temperature - short terms

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1. Introduction

AISI 441/DIN 1.4509 is a ferritic stainless steel stabilized with Nb and Ti and AISI 316L/DIN 1.4404 is an austenitic stainless steel. These alloys are often used for their strong oxidation resistance. They are widely studied at lower temperature for the nuclear industry, and studied consistently at 1100 °C for corrosion resistance during the hot rolling and annealing process [Saunders et al. (2008)]. However, most studies present variations from one to another because of the sensibility of these alloys depending on oxidation parameters. Indeed, temperature, water vapor and exposure time all have a strong impact on oxides scale. This study gives therefore a consistent look at the oxides generated in this range of parameters using complementary analysis techniques.

Regarded as essentially composed of Cr₂O₃ on the ferritic alloy, the thick oxide generated is actually more complex. An effort has been then made to identify elements, phases and their localization. If silicon oxides are known for a long time [Saunders et al. (2008)], Mn-spinels are often depicted differently [Brady et al. (2008)]. The oxides on the austenitic steel are also analysed in detail as variations have been observed depending on the water content in the atmosphere.

The study was originally aimed at a process which requires as a first step to obtain rapidly a thick and homogeneous oxide. It was therefore approached at first from the opposite aim of preventing corrosion. The results are, however, still of interest for annealing and hot rolling process investigations.

2. Experimental methods and materials

The composition of 441 and 316L stainless steels are shown in table 1. The average grain size is 40 μm for the ferritic steel and 17 μm for austenitic steel. Samples were received as skin-pass finish and cut in pieces of dimension 20x80x1 mm. Before treatment they were cleaned with acetone. The oxidation stage was performed in a belt furnace in which duration, atmosphere and temperature were set accordingly. The oxidizing atmosphere was produced from an argon flow of 0.3, 0.6 and 0.9 m³/h passing through a water tank set at 85 °C and then mixed with 3 m³/h of pure argon. 5%, 9% and 13% of humidity was therefore obtained. Raman micro spectroscopy has been operated with an Ar⁺ laser. The wavelength is 514 nm. XRD experiments are realized on a Philips X’Pert Pro MPD. X-Ray radiation is Cu-Kα. Grazing angles have been used to consider oxides rather than the bulk. 0.2° and 1° have been selected. EDS and SEM pictures were realized on a Philips XL30. XPS system is a Thermo-Fisher K-Alpha with an Al Kα source.

Table 1 – Chemical compositions (wt. %).

| Grades  | 1.4509 / 441 | 1.4404 / 316L |
|---------|-------------|--------------|
| C       | ≤ 0.02      | ≤ 0.03       |
| Si      | ≤ 1         | ≤ 1          |
| Mn      | ≤ 1         | ≤ 2          |
| Cr      | 17.5-18.5   | 17-19        |
| Mo      | -           | 2-2.5        |
| Ni      | -           | 10-13        |
| P       | ≤ 0.04      | ≤ 0.05       |
| S       | ≤ 0.03      | ≤ 0.03       |
| Ti + Nb | 0.6         | -            |
| N       | -           | ≤ 0.03       |
| Fe      | Bal.        | Bal.         |

3. Results and discussion

For 2.5 min of oxidation on 441 stainless steel, a very thin nano-scale layer of protective chromium oxide is formed. For all longer exposure times until 20 min, an almost homogeneous and much thicker layer, mainly consisting of Cr₂O₃, is produced. The thickness varies slightly and gradually from 4 to 20 min of oxidation (fig. 1) as well as from 5% to 13% of H₂O (fig. 2). The range of variation goes from 2 μm to 5 μm. In other terms, the kinetic of oxidation change drastically after 2.5 min of exposure time to reach a thicker oxide steadily developed. After this point, all samples present very similar oxides. Fig. 3 presents the Raman spectrum of the surface of the
sample and from the cross-section. Both areas are spotted in the optical view. There are Mn-Cr spinels mixed with the chromium oxide in the main layer (area 2). XRD results (fig. 4) indicate a decreased ratio of the spinel phase over corundum phase from 0.2° to 1° grazing angle. XPS and EDS (fig. 5) also confirm the presence of Fe and the increase of Mn in this outer part. Coupled with Raman results, the most external part of the oxides is thus a mix of Mn, Fe and Cr spinels. The presence of a layer of SiO₂ between the bulk and chromium oxide is also visible on fig. 5.

![Figure 1 - 441 alloy. Optical view of cross-sections. Influence of time with constant %H₂O.](image1)

![Figure 2: 441 alloy. Optical view of cross-sections. Influence of %H₂O with constant time.](image2)

![Figure 3: 441 alloy. Raman spectrum for sample Ar + 13% H₂O and 10 min. (1) top layer (2) main layer with corresponding optical view of the cross-section.](image3)
For all exposure times, 316L forms a thick oxide scale with iron oxides on its outer part and spinels for its inner part. Iron oxides are differentiated on fig. 6. For Ar + 5% H₂O, iron oxides for all durations are constituted of a single layer of wüstite Fe₁ₓO. For Ar + 9% H₂O and Ar + 13% H₂O, iron oxides are systematically constituted of a layer of hematite Fe₂O₃ over a layer of magnetite slightly enriched in chromium Fe₃₋ₓCrₓO₄. The lower oxidizing strength of the atmosphere for 5% H₂O may be the cause of this observation as the oxide state of Fe in FeO is lower. Due to these strong oxidation conditions, oxide scale is not always homogeneous and iron oxides spallation may occur. It is true in particular for longer oxidation time and higher water vapor content. The inner oxide, confirmed by Raman (fig. 6. area 3) and EDS is always composed of a mix of mainly Fe and Cr with additions of Ni in a spinel structure.
The amount of chromium in a stainless steel alloy is regarded as the main factor to increase the resistance to corrosion [Lacombe et al. 1993]. In these two alloys, this amount is very similar. Their oxide layers for the parameters studied here are however very distinct. For 316L steel, the mechanism involved is called ‘breakaway’. This configuration of oxides has been well studied but is not fully understood yet [Saunders et al. (2008)]. The very thin, nano-scale, chromium oxide existing at room temperature, breaks when the oxidation parameters become too strong in terms of temperature, duration or atmosphere. Several mechanisms are suggested to be involved [Saunders et al. (2008), Gheno et al. (2012), Saeki et al. (2012)]. The early layer formed is partially not adhesive and when the oxidizing molecules reach the chromium depleted bulk, the oxidation becomes steep. The significant augmentation of the diffusivity with temperature and the evaporation of chromium oxides are also considered responsible for the resulting oxides. But all of these reasons come from a lack of chromium reaching the surface to heal the defective former protective layer. Instead, it results in the use of iron as iron oxides and Fe-spinels, both present in the breakaway oxides. Yet, the main difference between austenitic and ferritic steels is the structure given by the presence or the absence of Ni. At high temperature, the diffusivity of chromium is higher in the ferritic lattice [Peraldi and Pint (2004)]. This difference of diffusivity is therefore regarded as the main cause of the difference of oxidation behavior between the two grades. When the initial protective layer is unable to prevent the oxidizing of the ferritic alloy, enough chromium reaches the surface for the parameters used in this study. Indeed, ferritic stainless steels can also have a breakaway oxidation mechanism usually for longer durations or weaker alloys [Brady et al. (2008), Gheno et al. (2012), Saeki et al. (2012), Peraldi and Pint (2004)]. Si and Mn have been found to be essential in this range of treatment durations to avoid the breakaway [Ledoux et al. (2013), Gonzales et al. (2008)]. The formation of Mn-spinels onto the external layer limits the chromium oxide evaporation and the SiO$_2$ layer acts as a diffusion barrier. However, their presence in the ferritic alloy and their role in the oxidation do not justify the difference with the 316L as they are present in both grades. The grain size does not justify the difference in behavior either as grain size in 441 alloy is bigger than in 316L alloy.

4. Conclusion

Ferritic stainless steel 441 has a better corrosion resistance for the parameters studied. After 2.5 min of oxidation, it always results in a few microns thick main layer of Cr$_2$O$_3$ mixed with (Mn,Cr)$_3$O$_4$. More spinel phase, with Fe, Mn and Cr are present on the outer part. A thin layer of SiO$_2$ forms a barrier with the bulk. 316L alloy presents in all cases a breakaway type oxide scale. The amount of water vapor plays a role in the nature of the iron oxides formed on the outer part. FeO is observed for lower water vapor partial pressure and both Fe$_2$O$_3$ and Fe$_3$O$_4$ are present for higher water vapor content. Inner oxides are constituted of Fe-Cr-Ni spinels. The higher diffusivity of chromium in the ferritic lattice plays a major role by modifying the oxidation mechanism and thus, the oxides scale. As a result
from the original purpose of this work stated in the introduction, both grades produce a thick oxide. 441 Alloy is however more homogeneous without spallation issues.

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