Oxidation study of alumina–forming alloys

S. Chevalier¹, C. Houngninou¹, S. Paris¹,², F. Bernard¹, E. Gaffet², Z.A. Munir³, J.P. Larpin¹, G. Borchardt⁴

¹Laboratoire de Recherches sur la Réactivité des Solides, UMR 5613 CNRS, Université de Bourgogne, 9 avenue Alain Savary, BP 47870, 21078 Dijon cedex, France.
²Laboratoire Métallurgies et Cultures, UMR 5060 CNRS/UTBM, Site de Sévenans, 90010 Belfort, France.
³Department of Chemical Engineering and Materials Sciences, University of California, Davis CA 95616, USA.
⁴Institut für Metallurgie, Technische Universität Clausthal, Robert Koch Strasse 42, 38678 Clausthal-Zellerfeld, Germany.

sebastien.chevalier@u-bourgogne.fr

Abstract

Fe–20Cr–5Al, pack–aluminised Fe–30Cr model alloys and FeAl intermetallics have been oxidised in air under atmospheric pressure and isothermal conditions. The high temperature efficiency of the thermally oxide scales grown on these three materials were compared. The $k_p$ values deduced from the parabolic plots of weight gain curves showed that $\alpha$–Al$_2$O$_3$ composed the oxide scale on all samples oxidised at $T>1000$ °C. For lower temperatures, transient alumina phases are observed, except for the Fe–20Cr–5Al alloys, for which only $\alpha$–Al$_2$O$_3$ was characterised whatever the temperature up to 800 °C. The addition of a reactive element (Y or Y$_2$O$_3$) drastically improved the high temperature oxidation resistance of the tested materials, since the oxide growth rate decreased and the alumina scale adherence strongly increased. The oxide morphologies and the X–ray diffraction patterns helped to compare the high temperature behaviours under oxidant atmospheres.
Keywords: alumina–forming alloys, intermetallics, pack aluminisation, Al₂O₃ scales, reactive element, oxidation mechanisms.

Introduction

The high temperature resistance of steels is based on their capacity to form homogeneous, adherent and dense protective oxide scales. Alumina–forming alloys are good candidates to resist to high temperature oxidation atmospheres because alumina scales, especially α-Al₂O₃, exhibit the expected properties [1]. Intermetallic compounds, in particular aluminides, have been developed more recently. They exhibit also excellent high temperature resistance because they form Al₂O₃ scales acting as diffusion barriers during the oxidation process at high temperature [2,3,4,5,6,7,8,9]. The major problem encountered by intermetallic compounds is their brittleness, especially at room temperature, which creates difficulties during their production [8,9]. Another possibility to elaborate aluminides consists in introducing Al at the steel surface using an Al deposition technique. The pack cementation process has been commonly used to cover iron–based alloys and to form aluminides on their surfaces [10,11,12,13,14,15,16,17,18].

The aim of this work is to compare the high temperature oxidation properties of Fe–20Cr–5Al, pack–aluminised Fe–30Cr model alloys and FeAl intermetallics. Isothermal oxidation tests in the temperature range from 800 to 1200°C have been performed to check their ability to form alumina scales with the expected protective properties.

The use of two stage oxidation experiments under ¹⁶O₂ and ¹⁸O₂, the observation of transmission electron microscope (TEM) cross–sections and the use of more classical analytical techniques, as scanning electron microscope (SEM) and X–ray diffraction (XRD), help to describe the high temperature oxide scale growth and to understand the differences observed between the tested samples.

A particular attention will be paid to the role played by Y or Y₂O₃ addition on the high temperature oxidation behaviour when added to Fe–20Cr–5Al, pack–aluminised Fe–30Cr model alloys and FeAl intermetallics.
Experimental

Fe–20Cr–5Al “model” steels were classically elaborated in laboratory conditions by melting route. They were called “model” alloys since they did not contain any minor elements to avoid the well known influence of alloying elements on the high temperature corrosion behaviour [19,20,21]. There was a low amount of sulfur (S>10ppm) in order to delete the sulfur effect [22,23,24,25,26,27], which was also susceptible to drastically influence the high temperature behaviour. Yttrium was introduced as an alloying element in Fe–20Cr–5Al steels to get Fe–20Cr–5Al–0.1Y alloys.

The intermetallic compounds were produced via Mechanically Activated Field Activated Pressure Assisted Synthesis (MAFAPAS) technique [28]. It consisted in applying a mechanical pressure together with an electric field on mechanically activated powders. The complete process was described elsewhere [28,29,30,31,32] and yielded to obtain FeAl materials with controlled densification and nanostructure. Indeed, the density of the so-prepared FeAl samples was over 99 % with an average crystallite size of 40–80 nm. Yttria was added in FeAl intermetallics as dispersed particles during the mechanical activation of powders to obtain FeAl–0.1Y2O3 (wt. %). The quantity of yttria was determined according to the literature in order to avoid any “overdoping” effect, generally observed when the reactive element content is too high in metallic materials [33]. The aluminisation of Fe–30Cr “model ” alloys was carried out using the pack cementation technique which involved a pack mixture made of 15 wt.% Al, 3 wt.% NH4Cl and 82 wt.% Al2O3 powders. The metallic substrate was embedded in the pack mixture for 5 hours at 1000 °C under argon atmosphere, to avoid oxidation during the process. The surface of the so-coated steels was covered with (Fe,Cr)3Al and (Fe,Cr)Al intermetallic phases. The aluminised zone was around 100 µm thick [34]. Yttria was introduced as a coating elaborated via the Metal–Organic Chemical Vapour Deposition (MOCVD) technique, which was completely described elsewhere [35]. An Y2O3 film of around 200 nm thick was
applied on the aluminised surface or on the Fe–30Cr substrate, before the pack–cementation process.

The samples were oxidised over the temperature range from 800 to 1200 °C in laboratory air under atmospheric pressure. Two stage oxidation experiments were performed at 1000 °C. The samples were firstly oxidised in $^{16}\text{O}_2$ under 200 mbar pressure. The oxidant atmosphere was then evacuated without cooling the samples and replaced by $^{18}\text{O}_2$ (isotopic purity $\approx 91\%$) under the same pressure. The oxygen isotope profiles were monitored by Secondary Neutral Mass Spectrometry (SNMS) to understand the alumina scale growth mechanism [36,37,38,39,40,41,42,43].

The corrosion products were characterised by scanning electron microscopy (SEM) with a field emission gun (FEG) coupled with an energy dispersive X-ray analyser (EDX). The thermally grown oxides were identified by X-ray diffraction (XRD), using the K$_{\alpha_1}$ copper radiation ($\lambda=0.154056$ nm).

**Results**

**Isothermal oxidation tests**

Figure 1 exhibits the oxidation kinetic curves of the Fe–20Cr–5Al steel in the temperature range from 800 to 1200 °C.
Figure 1: Isothermal oxidation kinetic curves for Fe–20Cr–5Al steels in air under atmospheric pressure.

The weight gain curves obey the parabolic rate law whatever the studied temperature. The parabolic constants were deduced from the slopes of the $\Delta m/S=f(t^{1/2})$ plots and are summarised in Table 1.

<table>
<thead>
<tr>
<th>$k_p$ (g².cm⁻⁴.s⁻¹)</th>
<th>Fe–20Cr–5Al</th>
<th>Aluminised Fe–30Cr</th>
<th>FeAl “nanostructured”</th>
</tr>
</thead>
<tbody>
<tr>
<td>800 °C</td>
<td>9.4 $10^{-16}$</td>
<td>5.4 $10^{-14}$</td>
<td>5.8 $10^{-14}$</td>
</tr>
<tr>
<td>900 °C</td>
<td></td>
<td>4.3 $10^{-13}$</td>
<td></td>
</tr>
<tr>
<td>1000 °C</td>
<td>2.5 $10^{-13}$</td>
<td>4.9 $10^{-13}$</td>
<td>1.2 $10^{-14}$</td>
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<tr>
<td>1050 °C</td>
<td>6.4 $10^{-13}$</td>
<td></td>
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</tr>
<tr>
<td>1080 °C</td>
<td></td>
<td>2.1 $10^{-12}$</td>
<td></td>
</tr>
<tr>
<td>1100 °C</td>
<td>1.5 $10^{-12}$</td>
<td></td>
<td>5.6 $10^{-13}$</td>
</tr>
<tr>
<td>1200 °C</td>
<td>2.2 $10^{-11}$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 1: Parabolic constants ($k_p$) for Fe–20Cr–5Al, aluminised Fe–30Cr and FeAl samples oxidised in air under atmospheric pressure.

Figure 2 shows the weight gain curves for the aluminised Fe–30Cr oxidised from 800 to 1080 °C. The weight gains increase with the temperature and the curves obey a parabolic law, as depicted in Table 1.

Figure 2: Isothermal oxidation kinetic curves for aluminised Fe–30Cr steels in air under atmospheric pressure.
Figure 3: Isothermal oxidation kinetic curves for FeAl intermetallics in air under atmospheric pressure.

The kinetic curves of the FeAl intermetallic compounds oxidised at 800, 1000 and 1100 °C are shown in Figure 3. The kinetic curve at 800 °C obeys a parabolic law from the beginning of the oxidation process. At 1000 and 1100 °C, a rapid increase of the weight gain per unit area is observed during the first hour of oxidation, followed by a classical parabolic shape. The corresponding k_p values are summarised in Table 1.

For all tested samples, the k_p values are close to each other. Note that the parabolic rate constant is lower for Fe–20Cr–5Al at 800 °C and for FeAl at 1000 and 1100 °C respectively.

The effect of yttrium or yttria was tested at 1000 °C. The weight gain curves (Figure 4) clearly evidence that the addition of the reactive element or the reactive element oxide decreases the oxidation rate and the weight gain during 100 h oxidation, compared to the undoped materials (Figure 1 to 3). The single exception concerns the aluminised sample for which the yttria coating applied after the pack cementation process and which exhibits faster oxidation kinetics and a higher weight gain, compared to the only aluminised sample (Figure 2).
Figure 4: Isothermal oxidation kinetic curves for Fe–20Cr–5Al–0.1Y, Y2O3–coated Fe–30Cr followed by aluminisation, Aluminised Fe–30Cr followed by Y2O3 coating and FeAl–0.1 Y2O3 intermetallics in air under atmospheric pressure at 1000 °C.

Oxide scale characterisation

The morphologies of the oxide scales formed on Fe–20Cr –5Al steels at 1000 °C and 1100°C are different. At 1000 °C, the oxide scale is largely convoluted with anchor points of the scale on the metallic substrate. Many spalled areas were observed all over the surface. At 1100 °C, the oxide surface remains flat with much spallation. The oxide scale fracture cross-section exhibits coarse grains with some needles on the oxide surface. For all tested temperatures (from 800 to 1200 °C), α–Al2O3 was identified by XRD.

After 100 h at 1000 °C, the aluminised specimens are covered with small oxide grains. At 1080 °C, the oxide scale is slightly convoluted and still composed of small oxide grains. The oxide scale is mainly composed of α–Al2O3 with transient alumina phases at T<1000 °C.

The FeAl intermetallic compound exhibits an “hairy” microstructure, since its surface is covered with thin needles after 100 h at 1000 °C in air. At 1100 °C, the oxide scale is composed of classical small
platelets. The XRD experiments clearly evidenced transient alumina phases at 1000 °C, whereas $\alpha$-Al$_2$O$_3$ was only detected at 1100 °C.

Two-stage oxidation experiments

The SNMS profiles of $^{16}$O and $^{18}$O are presented in Figure 5. The experiments were performed at 1000 °C for the three tested materials. The oxygen isotope profiles clearly indicate the presence of two major peaks of $^{18}$O for the three materials: one peak is located at the oxide surface and the second main $^{18}$O peak is situated within the pre-existing oxide scale close to the metal–oxide interface (Figure 5 a to c).
Figure 5: Oxygen isotope profiles after two-stage oxidation experiments at 1000 °C in $^{16}$O$_2$ and $^{18}$O$_2$, a) Fe-20Cr-5Al, b) aluminised Fe-30Cr and c) FeAl intermetallics.

Discussion

The Fe-20Cr-5Al steels, aluminised Fe-30Cr steels and FeAl intermetallic compounds oxidation curves obey the parabolic rate law. $k_p$ values deduced from the parabolic plots are comparable for $T>1000$ °C. At lower temperature, the Fe-20Cr-5Al steel exhibited a $k_p$ value two order of magnitude lower than the values determined for the aluminised steel and the intermetallics. Grabke et al. [44] exposed Arrhenius plots of $k_p$ together with the corresponding alumina phases formed on NiAl: $\alpha$-Al$_2$O$_3$ at $T>950$ °C, $\theta$-Al$_2$O$_3$ at $850<T<950$ °C and $\gamma$-Al$_2$O$_3$ at $T<850$ °C. The comparison of our results to the Grabke's Arrhenius plots evidences that $\alpha$-Al$_2$O$_3$ is formed on the Fe-20Cr-5Al steels all over the temperature range (from 800 to 1200 °C), that transient alumina phases ($\theta$ and $\gamma$) grow at low temperature for the aluminised samples and the intermetallic FeAl compounds, whereas $\alpha$-Al$_2$O$_3$ is the main thermally grown phase for $T>1000$ °C on these two last specimens.

These conclusions are confirmed by the observation of the oxide morphologies and above all by the XRD analyses, which identify transient alumina phases at low temperature for the aluminised steels and the FeAl specimens.

The three tested materials possess better high temperature behaviour with the presence of yttrium or yttria, verifying hence the beneficial effects of the so-called reactive elements [45]. Note that for the aluminised steels, the way to introduce the yttria has a major influence on its high temperature oxidation performance. When it is introduced after the aluminisation process, the yttria coating does not improve the oxidation resistance; the oxidation rate and the weight gain during 100 h oxidation tests are higher, compared to those of the undoped aluminised samples. This detrimental phenomenon was already observed [33,46,47] and attributed to an “overdoping” effect and bad
incorporation of the reactive element within the growing alumina scale, inhibiting then any beneficial effect of the reactive element.

The oxidation mechanisms were clarified at 1000 °C on the Fe-20Cr-5Al steels, the aluminised Fe-30Cr steels and the FeAl intermetallic compounds. Both inward diffusion of oxygen and outward diffusion of aluminium participate to the growth of the alumina scales. Two-stage oxidation experiments at lower temperature (800 °C for example) are actually in progress to understand the effect of the presence of transient alumina phases on the transport mechanism of the thermally grown alumina scales.

Conclusions

The oxidation behaviour of Fe-20Cr-5Al steels, pack cemented aluminised Fe-30Cr steels and FeAl intermetallic compounds were tested in air between 800 and 1200°C. Kinetic results and XRD characterisation evidenced that the alumina scales were composed of $\alpha$-Al$_2$O$_3$ at high temperature (T>1000 °C). Transient alumina phases ($\theta$-Al$_2$O$_3$ and $\gamma$-Al$_2$O$_3$) were identified at lower temperature (T<1000 °C) for the aluminised and the FeAl specimens. $\alpha$-Al$_2$O$_3$ was the main phase even at 800 °C for the Fe-20Cr-5Al steel.

The introduction of yttrium or yttria gave the well known reactive element effect, which consisted in decreasing the oxidation rate and improving the oxide scale adherence. For the aluminised samples, the beneficial effects were observed when the yttria coating was applied prior to the aluminisation.

Outward diffusion of aluminium and inward diffusion of oxygen constituted the main diffusive specie paths during the growth of the alumina scales at 1000 °C.

The comparison of the preliminary results clearly evidenced that pack aluminisation or the use of FeAl intermetallic could replace FeCrAl steels in order to form protective alumina scales, even if it is obviously
to early to relate the base material microstructure to the alumina scale growth mechanisms.

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References


