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KINETICS AND MECHANISM OF THE OXIDATION PROCESS OF TWO-COMPONENT Fe-Al ALLOYS

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This work presents results of studies of the oxidation process of two-component Fe-Al alloys containing I) up to 7.2% Al, and II) from 18 to 30% Al. Kinetic measurements were conducted using the isothermal gravimetric method in the range of temperature I) 1073-1223 K and II) 1073-1373 K respectively, for 50 hours. The methods used in studies of the mechanism of oxidation included: X-ray microanalysis, X-ray structural analysis, metallographic analysis and marker tests.

This work is a continuation of studies described in Reference [1], both papers giving a full picture of the process of oxidation of Fe-Al alloys containing up to 7.2% and 18-30% Al.

A mechanism for formation of scales on the investigated alloys is proposed on the basis of the obtained results.
KINETICS AND MECHANISM OF THE OXIDATION PROCESS OF TWO-COMPONENT Fe-Al ALLOYS

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I. INTRODUCTION

The problem of oxidation of two-component Fe-Al alloys has been an object of interest for the Department of General Chemistry of the Institute of Metallurgy at the Czestochowa Polytechnics. In this paper we present the results of studies on kinetics, morphological structure, and the chemical and phase composition of the product of oxidation. This work is a continuation of earlier studies on the kinetics and mechanism of the oxidation of two-component Fe-Al alloys. Results of the first part, which involved compositions with lower contents of aluminum, have been published already in Archiwum Hutnictwa [1]. This part extends the problem to higher ranges of temperatures and to higher contents of aluminum.

2. EXPERIMENTAL PART

Investigations were carried out on 7 types of two-component Fe-Al alloys, which were prepared from iron EO-4 (the contents of impurities did not exceed 0.16%) and aluminum class A-0 99.999% pure. The contents of the basic alloy component in particular alloys are given, among others, in Table 1. The process of melting the alloys, making the structure uniform and preparing samples for tests is described in [1]. Alloys were obtained with contents of aluminum up to 30% in which the contents of impurities did not exceed 0.18%.

* Numbers in margin indicate pagination in foreign text.
Figure 1. Parabolic course of the process of oxidation of two-component alloy containing 0.68% Al in the temperature range 923-1223 K (scales for temperatures 923, 998 and 1073 on the left side, and for 1148 and 1223 on the right side of graph)

Figure 2. Parabolic course of the oxidation process of two-component alloys 2, 3, 4 and in the temperature range 1148-1223 K (scale for alloys 3 and 4 on the left, and for 2 on the right side of graph); stop = alloy, stop = alloys
Figure 3. Parabolic course of the oxidation of alloy 5 containing 18.08% Al in the temperature range 1073-1373 K.

Figure 4. Parabolic course of the oxidation of alloy 6 containing 22.6% Al in the temperature range 1073-1373 K.
Studies of the oxidation rate

Kinetic studies were carried out on a thermogravimetric apparatus using the continuous isothermal gravimetric method. The rate of oxidation was measured during the 50-hour oxidation process at temperatures of 923, 998, 1073, 1148, 1223, 1298 and 1373 K in air atmosphere.

Errors in determining the parabolic reaction rate constant would arise from inaccuracies in directly measured quantities such as: time, mass and surface area of sample, and variations in temperature. The error resulting from inaccurate measurements of time, at the applied frequency of reacting every 10 minutes, was small and did not exceed 5 sec, and was 0.8% on the average. The accuracy of weighing did not exceed $\pm 5 \cdot 10^{-4}$ g, which at the total mass increases of about 0.1 g meant 1% error.

Dimensions of samples (disc of diameter 25 mm and average thickness 2.5 mm) were measured with an accuracy of about 0.01 mm; hence, the relative error of the measurement of surface of samples was 0.5%. The constancy of temperature was ensured by application of an automatic electronic compensator type EPP-120 with regulating attachment and continuous recording, of the degree of accuracy 0.5 and sensitivity 0.1% of the measuring range. It ensured the temperature stability within the limits $\pm 3$ degrees, which corresponds to the maximum 0.5% error.

An error caused by changes of the metallic core surface during the reaction was also taken into account, following the work [2]. On the basis of average values obtained from seven measurements, graphs were prepared (Figures 1, 2, 3, 4, 5) showing the dependence of the square of mass increase of sample per unit area on the oxidation time. After no longer than 10 hours, one can see already
Table 1. Parabolic constants of the oxidation rate of Fe-Al alloys [g^W/m^4. hour]

<table>
<thead>
<tr>
<th>Alloy number</th>
<th>% Al</th>
<th>Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>923K</td>
<td>998K</td>
</tr>
<tr>
<td>1</td>
<td>0.08</td>
<td>0.2</td>
</tr>
<tr>
<td>2</td>
<td>3.29</td>
<td>2.1</td>
</tr>
<tr>
<td>3</td>
<td>7.5</td>
<td>4.7</td>
</tr>
<tr>
<td>4</td>
<td>8.69</td>
<td>2.3</td>
</tr>
<tr>
<td>5</td>
<td>18.08</td>
<td>0.32</td>
</tr>
<tr>
<td>6</td>
<td>22.60</td>
<td>0.12</td>
</tr>
<tr>
<td>7</td>
<td>28.88</td>
<td>0.22</td>
</tr>
</tbody>
</table>

the grouping of the majority of measurement points on a straight line, indicating that the process follows the parabolic law of oxidation. This regularity is particularly clear for alloys with lower contents of aluminum (below 18%), and is less pronounced for alloys with a higher percentage of aluminum (above 22%), which is connected probably with a larger measurement error arising for lower increases of the sample mass.

From straight-line portions of curves, we determined by a graphical method the rate constants of the reaction (Table 1). The mathematical analysis of the curves was performed on a digital
computer taking into consideration only the points obtained after 10 hours of the process. We tried to determine the probability that the points were following a parabolic relation by putting into a computer the values of the change of mass $y$ and the duration of process $x$ to fit the equation $y = ax^b + c$. The obtained values of $b$ were in the range $0.5 \pm 0.02$. On the basis of these results we can accept, with probability 0.98, the hypothesis for the parabolic course of the process. Taking into account the mentioned values of the measurement errors, and the error connected with determining the parabolic constant [3], we assumed that the relative error did not exceed 10%.

**Chemical X-ray microanalysis**

A clarification of the oxidation process requires thorough studies of chemical and phase composition, and morphological structure of the product of oxidation. We determined local changes of chemical composition of the scale (burnout) and metallic base by using an electronic microprobe type IXA-3A. The X-ray microanalysis for the contents of iron and aluminum was performed on samples previously subjected to the oxidation process at the highest temperature for a given alloy. Two time periods of the oxidation process were
Figure 6. Distribution of the concentration of iron and aluminum in scale and surface layer of alloy 2 with initial contents 3.78% Al. Temperature of oxidation reaction 1223 K.

Alloys with low contents of aluminum up to 7.2%, oxidized for 50 hours at a temperature of 1223 K, are covered with a scale (burnout) composed of two distinct layers separated by a gap (shown for illustration in Figure 6). The inside layer of the scale shows a certain enrichment in aluminum in comparison with the metallic base, whereas there is a lack of this component in the thicker outside layer. Both in the first and the second layer, the dominant component is iron. In the outside part of the scale, one can notice a sublayer with reduced contents of iron. The surface-adjacent layer of metal shows a slight increase of the aluminum contents towards the metal-scale boundary. In the analyzed layer of metal, there are clear indications of an increase in the contents of aluminum with a concurrent reduction of the contents of iron, which could be connected with internal oxidation. The scale formed on the investigated alloys 18-30% aluminum is thin (up to 0.03 mm), forming one layer, and consisting nearly exclusively of aluminum oxides (Figures 7 and 8).
Figure 7. Distribution of Fe and Al in scale and surface layer of alloy 5 with initial content 18.08% Al. Oxid. temperature 1373 K.

Figure 8. Distribution of Fe and Al in scale and surface layer of alloy 6 with initial content 22.6% Al. Oxid. temperature 1373 K.
In rare cases, for instance in alloy 5, one could notice traced amounts of iron oxides in the scale.

**X-ray investigations**

We performed an X-ray analysis using the powder method in order to identify the phase composition of the products of oxidation. Samples set aside for these investigations were subjected to oxidation using temperature and time, as in the case of studies with electronic microprobe. The results are presented in Table 2 (the data for identification of particular phases were taken from Reference [4]). It was found that the X-ray studies fully confirmed the results obtained with a microprobe. The scale formed on alloys containing up to 7.2% aluminum is composed nearly exclusively of iron oxides; the outside layer consists of hematite Fe₂O₃, magnetite Fe₃O₄ and wustite FeO, and the internal layer is composed of Fe₃O₄, FeO and mixed spinels FeO·Al₂O₃, characteristic for alloys 2, 3, 4. The common characteristic feature for analyzed alloys of this group is the appearance of a wustite phase, in the case of alloy 1 — in the whole range of investigated temperatures, and in the case of remaining alloys — above the temperature 1073 K.

The scale formed on alloys containing 18-30% aluminum is composed of aluminum oxide, while iron oxides and spinels appear only in trace quantities. The dominant phase of Al₂O₃ in the scale varies depending on the oxidation temperature. At lower temperatures (up to 1223 K), the low-temperature type of γ - Al₂O₃ appears, whereas at temperatures above 1223 K the high-temperature type of α - Al₂O₃ is formed, which is in agreement with the literature data [5, 6, 7].

**Metallographic studies**

The morphological structure of the product of oxidation was studied using the microscopic metallographic analysis of a transverse polished side of the sample, made perpendicularly to the original surface of disc covered with scale. The analysis was supplemented
Table 2. Phase composition of oxide scales on Fe-Al alloys.

<table>
<thead>
<tr>
<th>Alloy No.</th>
<th>% Al</th>
<th>923 K</th>
<th>1073 K</th>
<th>1148 K</th>
<th>1223 K</th>
<th>1373 K</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>w</td>
<td>z</td>
<td>w</td>
<td>z</td>
<td>w</td>
</tr>
<tr>
<td>1</td>
<td>0.08</td>
<td>Fe₂O₃</td>
<td>FeO·Al₂O₃</td>
<td>Fe₂O₃</td>
<td>FeO</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>2</td>
<td>3.78</td>
<td>Fe₂O₃</td>
<td>FeO·Al₂O₃</td>
<td>Fe₂O₃</td>
<td>FeO</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>3</td>
<td>4.70</td>
<td>Fe₂O₃</td>
<td>FeO·Al₂O₃</td>
<td>Fe₂O₃</td>
<td>FeO</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>4</td>
<td>7.20</td>
<td>Fe₂O₃</td>
<td>FeO·Al₂O₃</td>
<td>Fe₂O₃</td>
<td>FeO</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>5</td>
<td>18.08</td>
<td>Fe₂O₃</td>
<td>FeO·Al₂O₃</td>
<td>Fe₂O₃</td>
<td>FeO</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>6</td>
<td>22.00</td>
<td>Fe₂O₃</td>
<td>FeO·Al₂O₃</td>
<td>Fe₂O₃</td>
<td>FeO</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>7</td>
<td>28.00</td>
<td>Fe₂O₃</td>
<td>FeO·Al₂O₃</td>
<td>Fe₂O₃</td>
<td>FeO</td>
<td>Fe₂O₃</td>
</tr>
</tbody>
</table>

w- internal layer of scale; * - phase composition of scale growth; z- external layer of scale; ** - phase composition of scale between outgrowths; sl - traces
with tests carried out by the marker method which gave some idea about the relative role of reagents in the process of mass transport through the layers of the reaction product. A platinum marker was deposited on the polished surface of metal by spreading it in the form of short sections, and subsequently by oxidizing this prepared sample in the horizontal position. This is a method that ensures the best results.

The scale formed on alloys containing up to 7.2% Al is well delineated and consists of two distinct layers (Figures 9 and 10). The marker is located in the external layer of the scale, somewhat above the boundary with the internal layer. This fact indicates that the external layer of the scale is formed as a result of diffusion of metal ions out from the core. At the same time, some oxide exclusions are visible in the metallic phase, evidencing the occurring phenomenon of internal oxidation (Figure 11).

The oxidation product found on the analyzed samples containing 18-30% Al is a deposited layer. Only at temperatures of 1373 K is there a clear, thin scale, which forms a weakly adhering uniform layer. Experiments carried out with the aid of platinum markers failed to provide sufficient information with regard to the direction of diffusion of reagents in the scale, since the thickness of the scale obtained in 50 hours is of the order of 10-20 μm; hence, is less than the marker diameter (about 50 μm). At higher temperatures (above 1373 K), when thickness of scale increases, the marker is destroyed [9, 10].

DISCUSSION OF RESULTS

The studies of the kinetics of the oxidation of two-component Fe-Al alloys, and investigations of the chemical and phase composition as well as structure of the scale, coupled with results of studies of the diffusion of reagents obtained by the marker method, provided information on the corrosion of these materials. On the basis of
Figure 9. Microphotograph of a transverse polished side obtained on alloy 1 containing 0.68% Al. Temperature of oxidation 1223 K. Magnification 55 X.

Figure 10. Microphotograph of a transverse polished side obtained on alloy 2 containing 3.78% Al. Temperature of oxidation 1223 K. Magnification 75 X.
results of kinetic studies of the oxidation process, conducted in the temperature range 923-1373 K, we can state that, at the most, after 10 hours the process follows a parabolic law, independently of the alloy composition and the reaction temperature. It means that the slowest step, determining the rate of reaction, is the diffusion of reagents in the solid phase of the product of reaction.

Figure 12 shows that the changes in reaction rates, as a function of the contents of aluminum in the investigated alloys, are partially similar for all temperatures. This similarity can be noticed clearly in the regions of two groups: alloys with low contents of aluminum - up to 7.2% and alloys with 18-30%. On the other hand, the composition of the alloy distinctly affects both the phase character (Table 2), morphological structure and thickness of the scale (Table 3). Because of these changes, we can divide alloys into two groups: alloys with low contents of aluminum up to 7.2% and alloys with higher contents of aluminum of 18-30%.

**FE-Al alloys containing up to 7.2% Al**

The dependence of the rate constant of the oxidation of Fe-Al alloys on the temperature is presented in semi-logarithmic form in Figure 12. The graphs show breaks which evidence the effect of temperature on the oxidation mechanism of the substance. In all the alloys containing up to 7.2% Al, one can see a distinct increase of the reaction rate as the temperature rises (above 1023 K for alloy 2, above 1073 K for alloy 3, above 1123 K for alloy 4). The authors think that this is connected with changes in the phase structure of the scale occurring at a temperature dependent on the contents of aluminum. As a result of these changes, the ratio of concentration of phases which form the composition of the internal layer of scale is a function of temperature for this group of materials. Namely, at temperatures of 1073 K and lower, the predominant phase in this layer is the spinel phase Fe(Fe$_{2-x}$Al$_x$)$_{10}$O$_4$, whereas at higher temperatures there appears also the wistite phase whose amount increases as the temperature rises (Table 2). In the light of kinetic studies and the
Figure 11. Microphotograph of a transverse polished side of near-surface layer of metallic core and scale, obtained on alloy 2 containing 3.78% Al. Temperature of the reaction of oxidation 1223 K. Magnification 350 X.

Figure 12. Dependence of the rate constant of oxidation reaction on the temperature for Fe-Al alloys.
results on the chemical and phase composition and morphological structure of the scale, the following model for the oxidation of these two-component alloys is proposed (Figure 13). At first, a tri-layer scale characteristic for pure iron grows on the surface of alloy, and at the same time oxidation of aluminum takes place in the metallic phase. Growth of the external scale occurs because of the out-of-core diffusion of ions and electrons of iron, whereas the internal region of oxidation appears as a result of the towards-the-core diffusion of oxygen through pores and crevices formed in the scale. The source of oxygen for the process of internal oxidation might as well be the dissociation of the scale. As thickness of the scale increases, its contact with the metallic core becomes interrupted in the second stage and a gap is formed. In this way, conditions are created for dissociation of the wistite layer and for secondary reactions of oxygen arising from decomposition of FeO with iron ions coming from the metallic phase. As a result, a porous internal layer of wistite is formed.

In the discussed cases, the velocity of diffusion governs the rate of growth of particular scale layers. Hence, one could conclude that the wistite layer because of its thickness plays the decisive role in the scale formation. The appearance of the wistite phase in the scale is characteristic for the analyzed alloys oxidized at temperatures above 1073 K.

Summing up the results pertaining to the kinetics and mechanism of the oxidation of alloys 2, 3 and 4, we can state that the alloying additive, Al, decisively affects the rate of reaction and chemical composition of the scale. This alloying additive improves the heat resistance of the alloy and causes the formation of a protective layer of spinel or oxide structure. The mobility of ions of aluminum and in iron in the spinel phase is slower than the mobility of iron ions in the wistite or even magnetite layer.
Fe-Al alloys containing 18-30% Al

Graphs of dependence of the rate constants of oxidation $k_p$ on temperature are shown in semi-logarithmic form in Figure 12. The curves characterizing the high-aluminum alloys (5, 6, 7) show two straight lines with different slopes. The fact that this dependence is not rectilinear throughout its entire course but undergoes a break (change of slope) could be caused for two reasons: either there are structural changes in the metallic core itself, or there is a change in the phase composition of the scale at temperatures close to the temperature corresponding to the observed "break". If we accept as true the first alternative, i.e., the appearance of structural changes in the alloy, then the break should be explained on the basis of the equilibrium diagram of the Fe-Al system. We found on the basis of the X-ray microanalysis that, during oxidation of high-aluminum alloys, there is a selective oxidation of aluminum, which leads to impoverishment of this component on the external surface of the metallic core (Figures 7, 8), which is in agreement with the literature data [11]. Such a reduction in the concentration of aluminum could cause changes in the structure of surface-adjacent layer of alloy. As the diagram of phase equilibria of the system Fe-Al shows, in the range of concentrations 18-30% Al there are two structures: one with ordering of the type Fe$_3$Al, and the other of the type FeAl. These structures are characteristic for temperatures lower than 973 K [1]. The mentioned change of structure could affect the shape of the curve only at temperatures lower than the analyzed ones. The observed break in the curve characterizing a change in the oxidation rate with change of temperature occurs in the temperature range 1223-1273K (Figure 12). Under these conditions the possibility of structural changes is not probable, and the explanation of the above "break" based on the assumption of structural changes in the alloy is problematic.
Table 3. Thickness of scale formed on surface of the oxidized Fe-Al alloys.

<table>
<thead>
<tr>
<th>Type of alloy</th>
<th>% Al</th>
<th>Heating temperature</th>
<th>Time, hours</th>
<th>Thickness of scale</th>
<th>Zone of Porous</th>
<th>External layer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Internal</td>
<td>External</td>
<td>Total</td>
</tr>
<tr>
<td>1</td>
<td>0,88</td>
<td>1223</td>
<td>60</td>
<td>0,2</td>
<td>1,1</td>
<td>1,3</td>
</tr>
<tr>
<td>2</td>
<td>3,09</td>
<td>1223</td>
<td>60</td>
<td>0,5</td>
<td>0,6</td>
<td>1,1</td>
</tr>
<tr>
<td>3</td>
<td>7,42</td>
<td>1223</td>
<td>50</td>
<td>0,05</td>
<td>0,05</td>
<td>0,1</td>
</tr>
<tr>
<td>4</td>
<td>8,69</td>
<td>1223</td>
<td>100</td>
<td>0,05</td>
<td>0,11</td>
<td>0,16</td>
</tr>
<tr>
<td>5</td>
<td>18,08</td>
<td>1373</td>
<td>240</td>
<td>—</td>
<td>—</td>
<td>0,03</td>
</tr>
<tr>
<td>6</td>
<td>22,0</td>
<td>1373</td>
<td>240</td>
<td>—</td>
<td>—</td>
<td>0,02</td>
</tr>
<tr>
<td>7</td>
<td>29,88</td>
<td>1373</td>
<td>240</td>
<td>—</td>
<td>—</td>
<td>0,01</td>
</tr>
</tbody>
</table>

Figure 13. Diagram of the mechanism II of the formation of heterophase multilayer scale on the analyzed two-component Fe-Al alloys containing up to 7.2% Al.
Figure 14. Mechanism for the formation of an oxide scale on the analyzed high-percent Fe-Al alloys, on the basis of the author's investigations (directions of diffusion of the reagents were taken on the basis of Reference /15/).
It remains, therefore, to consider the second assumption that changes in phase composition of the scale are responsible for the "break" occurring on the curve (Figure 12). A change of the phase composition of the scale may be caused by the appearance of new phases or by phase change of the reaction product [13, 14].

The X-ray structural studies of scale formed on analyzed alloys have shown (Table 2) that in a layer formed at the temperature 1373 K there are trace amounts of iron oxides in addition to the basic phase \( \alpha-Al_2O_3 \). But at the temperature 1073 K, the presence of iron oxides was not observed (Table 2), and the basic layer of the scale is formed by aluminum oxides at the phase \( \gamma-Al_2O_3 \).

The reason for a change of the oxidation rate with the rise of temperature may be, therefore, either the appearance of iron oxides at higher temperatures or a phase transition of aluminum oxides. It does not appear probable that iron oxides appearing in trace amounts could cause such a distinct change in the activation energy (Figure 12). It seems, therefore, highly probable that the appearance on the surface of the oxide \( \alpha-Al_2O_3 \) which has a higher activation energy than the oxide \( \gamma-Al_2O_3 \), is a factor accelerating the oxidation process.

On the basis of the results of these studies, supplemented by the literature data concerning the directions of diffusion of reagents [15], we offer the following model of the formation mechanism of the scale on alloys analyzed (Figure 14). In the first stage of reaction, crystalline embryos are formed, probably of both oxides - iron and aluminum. The possibility of the formation of aluminum oxide is, however, larger than of the hematite layer, since the absolute value of the free energy of formation of aluminum oxides is considerably greater than that of iron oxides. The contents of aluminum are considerable; hence in a short time the whole surface of metal becomes covered with a uniform dense layer of aluminum oxides. The formed layer of \( Al_2O_3 \), having nearly no defects, has considerable resistance to both the out-of-core diffusion of cations of two metals and to into-the-core diffusion of
gaseous oxygen [16]. For these reasons, the thickness of the scale increases only little under these conditions.

CONCLUSIONS

1. The oxidation of all the investigated Fe-Al alloys occurs after passing the initial stage (up to 10 hours), according to the parabolic kinetic law, indicating that the total oxidation process is governed by the process of diffusion. Kinetics of the oxidation process of alloys heated in air atmosphere depends on the temperature and composition of the alloy, the rate of reaction decreasing with the higher percentage contents of aluminum.

2. The dominant component of the scale on alloys containing up to 7.2% Al is iron, and on alloys containing 18-30% Al - aluminum. In the case of alloys with low contents of Al, we find also aluminum oxides in the metallic phase, formed in the process of internal oxidation. As the content of aluminum in the alloy increases, spinels appear in the scale, and with a still further increase of aluminum they turn into oxides - \( \gamma-Al_2O_3 \) and \( \alpha-Al_2O_3 \). A layer of spinels or aluminum oxides slows down the process of oxidation.

The authors wish to thank Prof. Dr. Teodor Werber from the Institute of Solid State Chemistry at AGH in Cracow for valuable remarks concerning this work and work described in References [1,3], and for his considerable help in metallographic studies.

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