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Oxidation by oxygen of liquid Al-Mg alloys during elaboration process

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Al-Mg ; oxidation ; oxygen pressure ; kinetics ; growth reactivity

Abstract
The oxidation into MgO of an industrial Al-Mg alloy in the liquid state has been studied at 700°C by thermogravimetry. Since the kinetic curves were not reproducible, it was not possible to use them directly. Thus, it has been necessary to use a method based on the isolation method to obtain the influence of oxygen partial pressure on the areic reactivity (mole of MgO.m⁻².s⁻¹) of growth of MgO.

Introduction
During the elaboration of aluminium-magnesium alloys, the surface of the liquid metal may be oxidised, mainly leading to the formation of MgO. The industrial system is quite complex, since several gases of the atmosphere can react with the alloys (oxygen, water vapour, nitrogen, carbon dioxide...).

So, we have studied a « simplified » system : the oxidation of an Al-Mg 5% alloy by oxygen. The aim of this work is to obtain the influence of the oxygen pressure on the areic reactivity (mole of MgO.m⁻².s⁻¹) of growth of MgO, and to propose a reaction mechanism able to account for the experimental results, in order to predict the behaviour of the alloy in industrial conditions of elaboration and to limit its oxidation.

After giving the oxidation kinetic curves, the methodology used for the kinetic study of the MgO growth will be detailed. Finally, the influence of the oxygen pressure on the areic reactivity of growth of MgO will be obtained, and thanks to the results of an IR study, some indications for a reaction mechanism will be given.

Experimental
The alloy is an industrial Al-Mg 5% alloy, supplied by Pechiney. The samples are cylinders of 1mm height and 9mm diameter (this height was chosen in order to ensure an constant oxygen pressure around the sample). For higher samples (15mm to 3.5mm), more oxidised parts were observed (a ring of white powder at the surface of the samples), the localisation of this ring along the cylinders varying with the oxygen pressure fixed for the experiment. Thus, we have assumed that a gradient of oxygen pressure exist along the high samples. This was not observed for the 1mm-high cylinders.

The oxidation of the liquid alloy was followed by isothermal thermogravimetry at 700°C (thermobalance SETARAM TAG 24). The experiments were carried out at atmospheric pressure, under a flowing mixture of helium and oxygen (1l.h⁻¹), the partial pressures being controlled by mass-flowmeters (Brooks 5850E).

The heating and the melting of the samples were carried out under oxygen at atmospheric pressure since it has been observed that a high oxygen pressure allows to limit the oxidation of the liquid alloy; the oxygen pressure chosen for the experiment is established after 30 minutes of isotherm at 700°C. The experiments of simultaneous calorimetry and
thermogravimetry were performed using a SETARAM TG-DSC 111. In situ IR study was carried out with a Spectra Tech cell (model 0030-13) and a spectrometer Biorad FTS 185. Micrographs were obtained on a scanning electron microscope Jeol JSM 840.

**Experimental results**

As shown in a previous study [1], MgO is the first phase which appears during the reaction, up to 2% weight increase. Above 2% (that is a residual magnesium activity in the alloy lower than 0.023), MgAl₂O₄ is the thermodynamically stable phase as represented in figure 1, its formation being confirmed by X-Ray diffraction. The kinetic curves will then be limited to a weight increase of 2%.

![Thermodynamic stability of MgO, MgAl₂O₄ and Al₂O₃ phases. Oxygen pressure at equilibrium versus magnesium activity.](image)

During the oxidation experiments, a thin solid MgO layer is formed when the temperature increases, and when the alloy melts, it remains inside this solid layer which envelops the liquid metal. This layer undergoes strong deformations during the experiments, it is very rough and irregular. Its aspect varies from one experiment to another, as shown by the SEM micrographs of figure 2, which represent the surface of two samples oxidised up to the same weight uptake (0.7%), in the same conditions of temperature and pressure (P₀₂=400hPa).

![Surface morphology of two samples oxidised up to a weight uptake of 0.7% (T=700°C, P₀₂ = 400 hPa).](image)
SEM observations have also shown that the oxide layer is grey and formed of small MgO grains (about 0.5 μm) (figure 3a); moreover, some holes are observed inside the samples, in which MgO cubes are located (5 to 10 μm) (figure 3b). The shape of the curves obtained for oxygen pressures ranging from 200hPa to 800hPa is given in figure 4: the curves are very irregular and not reproducible, which is not surprising since the samples undergo very important morphological changes during the reaction.

![Image of SEM observations](image1)

**Figure 3: Morphology of the MgO oxide issued from the oxidation of Al-Mg5% alloy: surface of the samples (a) and in the holes (b).**

![Graph of weight uptake](image2)

**Figure 4: Curves of weight uptake and non reproducibility (T=700°C, P_{o2}=400hPa)**

**KINETIC STUDY OF THE OXIDATION OF THE ALLOY**

1: Method.

The transformation of a solid involves the processes of nucleation and growth of the new phase (MgO in our case). The modelling of the transformation is simplified when the following assumptions are made (which can be verified experimentally):

(i) the system is in a quasi-steady state

(ii) the derivative of the fractional conversion α versus time, which be also called the rate, can be written:

\[ R = \frac{d\alpha}{dt} = \Phi \cdot E \]  \[ \text{[E.1]} \]
where Φ is the areic « growth reactivity » of MgO (in mol.s⁻¹.m⁻²), which depends on the physico-chemical variables (pressure P, temperature T, magnesium activity...), and E is the « space function » (m².mol⁻¹), characteristic of the extent of the reaction area where the rate-limiting step of the transformation occurs. E depends on the time and on the history of the solid from the beginning of the transformation up to the considered instant [2]. When these assumptions are verified, an experimental method, based on the isolation method [3], can be used to obtain directly the variations of Φ with the physico-chemical variables (particularly the oxygen pressure).

2: Results.

2.a: Quasi-steady state

It can be shown [4] that when a system is in a quasi-steady state, the curves giving the reaction rate versus time, obtained with two different experimental techniques, are affine, the axis being the rate axis and the direction the time axis. We have chosen to measure simultaneously the oxidation rate by thermogravimetry and calorimetry (heat flow). The curves of rate of weight uptake and heat flow versus time have to be superimposed in two axis systems with different ordinate scales. Figure 5 shows the curves obtained at 700°C for an oxygen pressure of 200hPa.

Figure 5: Quasi-steady state test (700°C, P_O2=200 hPa)

During the temperature increase (10°C/min), the alloy melting (at about 648°C) induces a strong endothermic signal, which hides the exothermic signal due to the oxidation up to about 80 minutes (or Δm=0.4%). Then, the curves are superimposed. Thus, the approximation of the quasi-steady state is valid at least from a weight uptake of 0.4%, i.e. a fractional conversion equal to 0.12.

2.b: « Φ.E » test.

In order to verify that the rate R can be written as in Eq.[E.1], we use the isolation method, which consists into changing suddenly a physico-chemical variable (P, T) from a value Y₀ to a value Y₁, at a given fractional conversion. Practically, we have changed the oxygen pressure from 200hPa to 400hPa, at various fractional conversion, as indicated on figure 6. Let Rᵦ₀(P₀, αᵢ) and Rᵦ₁(P₁, αᵢ) be the rates before and after the pressure change at the fractional conversion αᵢ, they can be written (cf. [E.1]) :

\[ Rᵦ₀(P₀, αᵢ) = Φ(P₀) E(αᵢ) \]
\[ Rᵦ₁(P₁, αᵢ) = Φ(P₁) E(αᵢ) \]

so, their ratio is:
\[ \frac{R_a(P_1, \alpha)}{R_b(P_0, \alpha)} = \frac{\Phi(P_1)E(\alpha)}{\Phi(P_0)E(\alpha)} = \frac{\Phi(P_1)}{\Phi(P_0)} \]  \[ \text{E.2} \]

The ratio of the rates before and after the pressure change does not depend on the fractional conversion: thus, the « \( \Phi \cdot E \) » test is verified if this ratio is constant, whatever the fractional conversion at which the pressure is changed is.

![Fractional conversion vs time](image)

*Figure 6: application of the isolation method to the « \( \Phi \cdot E \) » test: oxygen pressure changes at various fractional conversions.*

The test results are indicated in figure 7: two domains of weight uptake can be observed for which the « \( \Phi \cdot E \) » test is verified: the rates ratio is constant for a fractional conversion between 0.09 and 0.21 (weight uptake between 0.25% and 0.6%), and then between 0.3 and 0.6 (weight uptake between 1 and 2%), but the ratio value is different in each domain.

![Fractional conversion vs ratio](image)

*Figure 7: « \( \Phi \cdot E \) » test by changing the oxygen pressure from 400hPa to 200hPa.*

Besides, it has been verified that when the pressure change is carried out at a given fractional conversion, the value of the rates ratio obtained in various experiments is nearly the same (error less than 10%). Thus, the reactivity \( \Phi \) is reproducible, the non reproducibility of the kinetic curves (figure 4) comes from the variations of the space function \( E \).
3: Variations of $\Phi$ with $P_{O_2}$ in each domain of weight uptake.

The experimental method to obtain the variations of $\Phi$ with $P_{O_2}$ is also based on the isolation method: in that case, several changes are carried out at a given fractional conversion $\alpha$, from an oxygen pressure $P_0$ (here 200hPa) to various pressures $P_i$ (the temperature being constant).

The ratio of the rates before and after the change, equal to $\frac{\Phi(P_i)}{\Phi(P_0)}$ from [E.2], is proportional to the variations of $\Phi$ with $P_{O_2}$, ($\Phi(P_0)$ being a constant).

The variations of $\Phi$ with $P_{O_2}$ were obtained by changing the oxygen pressure at a fractional conversion 0.21 in the first domain of weight uptake ($\Delta m<0.7\%$) and at 0.39 in the second domain of weight uptake ($1\%<\Delta m<2\%$). The figure 8 shows that in each domain, the growth reactivity decreases when the oxygen pressure increases, which is quite unusual for oxidation reactions.

![Graph showing variations of $\Phi$ with oxygen pressure in each domain of weight uptake](image)

*Figure 8: Variations of the growth reactivity $\Phi$ with oxygen pressure in each domain of weight uptake.*

5. Discussion and conclusions

We have verified that we could assume that the system is in a quasi-steady state, and that the oxidation rate could be written as a product « $\Phi \cdot E$ ». These results are important for a further modelling of the reaction (with a reaction mechanism in elementary steps), because they mean that it will be possible to use the assumption of the rate-limiting step to calculate the rate laws issued from the reaction mechanism. Comparing these laws to the experimental variations of $\Phi$ with $P_{O_2}$ (Figure 8) will normally lead us to determine the rate-limiting step of the reaction and thus the appropriate rate law for $\Phi$. However, a qualitative description of the reaction mechanism can be attempted.

Usually, the mechanisms proposed to describe the oxidation of metals lead to an areic growth reactivity $\Phi$ which increases when the oxygen pressure increases [1].

In our case, to account for the inhibiting effect of oxygen on the growth reactivity of MgO, we can consider a reaction mechanism involving two kinds of parallel oxygen adsorptions on the MgO surface, since several oxygen adsorbed species have already been observed on MgO [5]. Moreover we have carried out in-situ IR experiments during the oxidation of solid magnesium, showing that the MgO formation is accompanied by the appearance of a band at 1016 cm$^{-1}$, characteristic of O$_2^-$ species, according to [6]. The IR spectra are given in figure 9.
These O$_2^-$ species are very stable, since the band at 1016 cm$^{-1}$ is not modified at 500°C even when oxygen is suppressed from the gaseous atmosphere. They are supposed to be inactive for the oxidation, and to occupy adsorption sites. When the oxygen pressure increases, their concentration at the MgO surface increases as well, which could explain the experimental variations of $\Phi$ with the oxygen pressure.

Moreover, the proposal of a mechanism may lead to an interpretation for the existence of the two domains of weight uptake, in which $\Phi$ is not the same. These domains could be due for example, to a change in the MgO growth mechanism or to a change in the rate-limiting step of the same mechanism. Another explanation could be the predominant formation of the small grains in one part and of the larger cubes in the other part.

References


Figure 9: In-situ IR spectra during recorded during the oxidation of solid magnesium at 500°C by oxygen.