Some kinetics aspects of chlorine-solids reactions

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Abstract

The present paper describes detailed kinetics investigations on some selected chlorine-solid reactions through thermogravimetric measurements. The solids studied in this article include chemical pure oxides and sulfides as well as their natural bearing materials. The chlorinating agents employed are gaseous mixtures of Cl₂+N₂ (chlorination), Cl₂+O₂ (oxychlorination), and Cl₂+CO (carbochlorination). Results are presented as effects of various parameters on the reaction rate of these solids with these chlorinating agents. It was observed that the reactivity of these solids towards different chlorinating agents varied widely. Sulfides could be chlorinated at room temperature, while carbochlorination of chromium (III) oxide was possible only above 500 °C. The variation of the chlorination rate of these complex materials with respect to gas velocity, composition and temperature enabled us to focus some light on the plausible reaction mechanisms and stoichiometries. The obtained results were used for selective removal of iron from chromite concentrates, extraction of valuable metals from sulfide materials, purification of MgO samples, etc.

Keywords

Thermogravimetric analysis; Complex oxides; Sulfides; Chlorination; Oxychlorination; Carbochlorination; Kinetics.

1. INTRODUCTION

It is well known that chlorine possesses a high reactivity towards different substances at low temperatures. Further, many metal chlorides and/or oxychlorides have low boiling points and also have appreciably different vapor pressures at a particular temperature. These aspects led the “chlorine technique” as the subject of much attention in areas like mineral beneficiation, metal extraction and waste treatment. Many important articles devoted to the chlorination can be found in the literature[1]. The present paper mainly focuses on the kinetics aspects of chlorine-solids reactions approached through thermogravimetric measurements. Besides, higher selectivity of the chlorination methods by
employing different chlorinating medium such as oxychlorination (Cl₂+O₂) or carbochlorination (Cl₂+CO) is also discussed. Below paragraphs depict some examples where detailed kinetic studies were made.

The first example presents how Cl₂ can be used at low temperature for extraction of non-ferrous metals from sulfide concentrates (CuFeS₂, PbS, ZnS, FeS₂, etc.). It was observed that these sulfides interact with chlorine even at room temperature\(^{2-4}\). At about 300 °C, generated iron and sulfur chlorides were volatilized whilst the valuable metal chlorides (CuCl₂, PbCl₂, and ZnCl₂) were essentially concentrated in the chlorination residues. More than 95% of copper contained in the concentrate was recovered in the residue at 300 °C as water soluble CuCl₂.

The second case study presented is the upgrade of lean chromite ores and/or concentrate. It may be mentioned that, physical beneficiation of this material is almost impossible. Chlorination of chromite was carried out in presence of a reducing agent (for example CO). Results obtained earlier\(^{2, 6 y 7}\) indicated that the carbochlorination (Cl₂+CO) of this material at about 550 °C led to preferential extraction of iron. The chlorination extent of chromium was negligible and the Cr/Fe ratio was almost doubled in the residues, compared with that of the raw sample. Carbochlorination at higher temperatures led to a total extraction of chrome elements as chlorides (CrCl₃, FeCl₃, AlCl₃, and MgCl₂). A selective condensation of the gaseous phase allowed the recovery of almost pure MgCl₂. Similarly, the chromium chloride was separated from those of iron and aluminum.

The third example presents the selectivity of chlorination reaction in the presence of oxygen. This approach was used for the selective chlorination of elements contained in the chromite\(^{12-15}\) and for purification of magnesium oxide through selective chlorination and removal of iron oxide\(^{16}\).

In order to have better understanding of the chlorination (Cl₂), carbochlorination (Cl₂+CO) and oxychlorination (Cl₂+O₂) behavior of various raw materials, and detailed kinetic studies of these solids with various chlorinating gas mixtures were undertaken. The main objective of this article is to demonstrate the usefulness of simple thermogravimetric analyses (TGA) methods for effective measurement of chlorination kinetics of above mentioned selected solid samples with various chlorinating gas mixtures. The obtained kinetic information can be subsequently used for the upgrade of lean ores, such as chromites, complex sulfides, etc., and/or extraction of several valuable metals such as Cu, Mg, Pb etc.

### 2. METHODOLOGY OF THE KINETICS STUDY

The methodology adopted for the kinetics studies of various solid samples with different chlorinating gaseous mixtures consists in the following:

1. thermogravimetric analysis (TGA) using non-isothermal conditions,
2. batch-boat experiments and analysis of the solid products (SEM, microprobe, XRD, chemical analysis, etc.),
3. comparison of the reaction rate with that of volatilization of the reaction products,
4. determination of adequate gaseous mixture flow rate,
5. effect of the reactive gases proportions on the reaction rate,
6. determination of the apparent reaction orders with respect to the reactive gases,
7. evaluation of the apparent activation energy of the reaction (effect of temperature),
8. mathematical data processing in order to define the geometry of the reaction interface,
9. indication of the possible rate controlling step.

Detailed results of the chlorination kinetics of different solids were reported in literature\(^{12-15}\). Only some typical examples will be described illustrating the stages of these kinetics studies. The materials and experimental procedure are briefly depicted in the next section. However, more accurate conditions for each kinetics example will be given during the development of each individual kinetic study.

### 3. MATERIALS AND EXPERIMENTAL PROCEDURE

The sulfides' samples used in this investigation were the chalcopyrite (CuFeS₂), pyrite (FeS₂), galena (PbS) and sphalerite (ZnS). The elemental and mineralogical compositions of these samples of natural origin were determined by Scanning Electron Microscopy (SEM) and X-ray diffraction (XRD) analyses. Besides, a high grade copper concentrate "HGCC" was also used. It contains about 81.7% CuFeS₂, 6.4% ZnS, 5.7% lead compounds, 3.0% iron sulfides and less than 1.0% SiO₂. The oxides' samples of chromium (Cr₂O₃), iron (Fe₂O₃) and magnesium (MgO) were supplied by PROLABO (Paris, France). They are of synthetic origin and have a purity as high as 99%. The chromite sample used for this study was obtained from a chromite concentrate by successive separations in dense liquor in order to separate the gangue from the chromite.
mineral. Results of the microprobe and classical chemical analysis suggest that the chromite sample could be represented as: $(\text{Fe}^{2+}_{0.30}, \text{Mg}_{0.70})(\text{Cr}_{1.56}, \text{Al}_{0.37}, \text{Fe}^{3+}_{0.07})\text{O}_4$. All used samples were in powder state with low specific surface areas indicating a week grains’ porosity.

The thermogravimetric analysis (TGA) were performed using a micro-electronic balance CAHN type C 1000. The accuracy of this type of apparatus is about 0.1% for a sensitivity of 0.5 µg. The $(\text{Cl}_2, \text{CO}, \text{O}_2, \text{N}_2)$ gas flows are measured separately. They can be mixed and homogenized. Almost all the residual moisture of these gases is eliminated by circulation through CaCl$_2$ columns. The gas inlet system is built so that the gases are preheated before reacting with the sample. The temperature and the weight of the sample are recorded continuously as a function of time by a data acquisition system.

The thermogravimetric analysis is run using continuous temperature raise or isotherm modes. The experimental protocol was the following:

1. an amount of sample of about 40 mg was scattered uniformly in a large quartz nacelle,
2. during the non isothermal treatment, the pure gases or gas mixtures ($\text{Cl}_2$, $\text{CO}$, $\text{Cl}_2+\text{N}_2$, $\text{Cl}_2+\text{CO}$, $\text{Cl}_2+\text{CO}+\text{N}_2$, $\text{Cl}_2+\text{O}_2$ and/or $\text{Cl}_2+\text{O}_2+\text{N}_2$) were flown through the reactor and at the same time the furnace power was switched on,
3. for the isotherm mode experiment the sample was preheated in a low nitrogen flow up to the desired temperature, this before flowing the reactive gases through the reactor,
4. at the end of the predetermined treatment time, the reactive gases were replaced by nitrogen and the furnace is lowered down, and the reactor cools down as a result of $\text{N}_2$ flow,
5. the nacelle was taken out from the reactor and weighted. The treatment product was set aside for different analyses.

Few batch-boat chlorination tests were also performed in a horizontal experimental set-up. This set-up was composed of a gas metering unit followed by a gas purification system and a horizontal furnace. The gaseous reaction products were recovered through condensers and the outlet gases were purified before their release to the atmosphere. The reaction products (chlorination residues as well as condensates) were examined by SEM, XRD and chemical analyses.

Data obtained from different treatments were initially plotted as evolution of % weight change as a function of time. All negative values represent % weight losses (% WL), whilst positive ones express the % weight gains.

4. RESULTS AND DISCUSSION

4.1. Thermogravimetric analysis using non-isothermal conditions

These experiments could give preliminary information on the temperature range for the kinetic studies. In other words, TGA allows rapidly the evaluation of the reactivity of various solids towards chlorinating gas mixtures. For example, in figure 1 the percentage weight change of several sulfides with respect to temperature during their chlorination with $\text{Cl}_2+\text{N}_2$ gas mixture under non-isothermal conditions with a heating rate (dT/dt) of about 20 °C/min are presented.

Several important information are obtained from this figure such as: chalcopyrite starts to react with chlorine even at room temperature, at higher temperature this reaction leads to the formation of cupric, ferric, and sulfur chlorides as final reaction products; for pyrite initial weight gain was observed due to the substitution of sulfur by chlorine, while the weight losses of this sample at temperature higher than 250 °C were attributed to the volatilization of ferric chloride; sphalerite and the galena started to react with chlorine at temperatures higher than 200 °C generating their respective chlorides. Full reaction and volatilization of the chlorination products of the studied sulfides took place at higher temperatures.

However, the chlorination of sulfides is very complex and is difficult to be explained only by their treatment in non isothermal conditions. In order to have an insight about the pure sulfides’ chlorination, a series of batch-boat tests was performed between room temperature and 750 °C for a reaction time of 2 h. Results were drawn as
evolution of weight change as function of the temperature in figure 2. A high grade copper concentrate “HGCC” composed essentially of chalcopyrite was also treated in same conditions. The chlorination residues and condensates were systematically analyzed by SEM and XRD analysis. The chlorination residues of HGCC were subjected to chemical analysis for copper and iron contents.

The presence of chlorine, as observed by SEM, in the chlorination residue obtained at room temperature for CuFeS$_2$, FeS$_2$ and HGCC indicates that chlorine had reacted with these sulfides. The peak intensity of sulfur decreased with the temperature rise, while that of chlorine increased. At 300 °C, SEM analysis showed that the residues obtained from chlorination CuFeS$_2$ and HGCC were sulfur-free indicating that full chlorination of the sulfides was achieved. XRD results showed the presence of CuCl$_2$, besides non reacted CuFeS$_2$, in the chlorination residue of chalcopyrite and HGCC samples from room temperature.

Condensates of chlorination of FeS$_2$, CuFeS$_2$ and HGCC were significantly observed at temperatures higher than 200 °C. These condensates were essentially composed of Fe and Cl for a chlorination temperature lower than 300 °C showing that iron was removed as FeCl$_3$ which is characterized by a vapor pressure of about 1 atm at temperatures close to 300 °C (Fig. 3). At temperatures equal and higher than 400 °C, copper...
was also detected in the condensate of CuFeS₂ and HGCC chlorination. Sulfur was not detected in the solid condensates regardless of the chlorination temperature, indicating that sulfur was essentially removed as SₓClᵧ (S₂Cl₂+SCl₂ and intermediates) which have sufficiently high vapor pressures (Fig. 3) to be transported out by the exhaust gases. The formation of sulfur dichloride is probably due to high partial pressure of chlorine in the chlorination process. The formation of cuprous chloride (CuCl) during chlorination of CuFeS₂ and HGCC was confirmed by XRD at temperature equal or higher than 600 °C. It is probably product of cupric chloride decomposition in spite of high partial pressure of chlorine.

The qualitative analyses such as SEM and XRD indicated that, in our conditions, the temperatures close to 300 °C were adequate for the full chlorination of pyrite and chalcopyrite and for the volatilization of sulfur and iron chlorinated compounds. The chemical analysis of chlorination residue of HGCC showed that complete iron extraction was achieved at 300 °C and all copper is concentrated in the residue. The theoretical limit of selective chlorination of HGCC (LScCh in figure 2) is fully consistent with % WL of sample and extraction extent of iron obtained at 300 °C. The copper extraction started at higher and about 40 pct of copper were extracted at 600 °C.

The sulfides of zinc and lead reacted with chlorine at temperatures equal to or higher than 150 °C leading to their respective chlorides. However, complete reaction of ZnS was achieved at temperature lower than that of PbS. The chlorination of these sulfides is also complicated by the formation of liquid phases (M.p. of ZnCl₂ is 283 °C) and/or formation of an eutectic at 450 °C in the PbS-PbCl₂ system.

The complete chlorination of the selected sulfides can be classified by following descending order FeS₂ > CuFeS₂ > ZnS > PbS. It seems that this sequence follows the evolution of vapor pressure of chlorides as shown by figure 3. Nevertheless, other phenomena could occur during chlorination of sulfide at high temperatures such as decomposition of chalcopyrite and pyrite to cubanite and pyrrhotite, respectively and/or volatilization of lead sulfide.

Pertinently to these results, the chlorination kinetics of these sulfides at temperature ≤ 300 °C were studied in details in former works[2-4].

4.2. Comparison of the reaction rate with that of volatilization of the reaction products

The reactions of chlorine with a solid generate chlorides which have particular vapor pressure at a given temperature. Consequently, temperature plays a crucial role in kinetic measurements, which defines the proportion of chlorides that volatilized to that of which remains in the reaction zone (unvolatilized). Use of TGA technique to measure the evolution of the reaction extent require that either the volatilization rate of the reaction products is higher than that of its rate of formation or the chlorinated products remain completely in the reaction zone under the studied temperature range.

Figure 4 illustrates an example of Cr₂O₃ carbochlorination using non-isothermal[2-10]. The reaction of Cr₂O₃ with Cl₂+CO can be described by equation (1).

\[
\frac{1}{3} \text{Cr}_2\text{O}_3 + \text{Cl}_2 + \text{CO} \rightarrow \frac{2}{3} \text{CrCl}_3 + \text{CO}_2
\]  

(1)

Assuming that chromium (III) chloride (CrCl₃) is the only chlorinated product of the reaction, the volatilization of CrCl₃ was tested in the same gaseous mixture (Cl₂+CO). Results are also depicted in figure 4. It is clear from this figure, that the rate of volatilization of CrCl₃ at temperatures above 500 °C is higher than its rate of formation through the reaction of Cr₂O₃ with Cl₂+CO. In these conditions, the observed weight loss of Cr₂O₃ reaction with Cl₂+CO could be a direct measure of the extent of Cr₂O₃ carbochlorination reaction.

The chlorination of the magnesium oxide represents another case study. The reaction of MgO with chlorine (Eq. (2)), in absence of a reducing atmosphere, takes

![Figure 4. TGA of Cr₂O₃ and CrCl₃ in Cl₂+CO.](image_url)

Figura 4. ATG del Cr₂O₃ y CrCl₃ en atmósfera de Cl₂+CO.
place at \( T > 600 \, ^\circ\text{C} \) \cite{2-10}. Magnesium chloride has melting and boiling points of 714 \, ^\circ\text{C} and 1,412 \, ^\circ\text{C}, respectively. One possible way to decrease the reaction rate of MgO with chlorine is to increase the partial pressure of oxygen in the system. This can lead to a volatilization rate of MgCl\(_2\), obtained according equation (2), higher than that of its formation.

\[
\text{MgO} + \text{Cl}_2 \rightarrow \text{MgCl}_2 + \frac{1}{2} \text{O}_2 \tag{2}
\]

A series of MgO chlorination tests was performed at 950 \, ^\circ\text{C}, using a Cl\(_2\)+O\(_2\) gas mixture with an oxygen content that varied from 0 to 66.7 \% (molar \%). Figure 5 shows the evolution of \% weight change of MgO sample versus time during its oxychlorination for the conditions mentioned above. This figure suggests that the chlorinating gas mixtures containing more than 90 \% of Cl\(_2\) lead to a chlorination rate that is higher than that of MgCl\(_2\) volatilization. For this reason, the kinetics of MgO oxychlorination was studied using a Cl\(_2\)+O\(_2\) gas mixture containing a chlorine content less than or equal to 80 \% \cite{8}. Then, the carbochlorination kinetics of MgO in Cl\(_2\)+CO was investigated at temperatures lower than the melting point of MgCl\(_2\) \cite{10} assuming that the MgCl\(_2\) volatilization is negligible.

### 4.3. Determination of adequate gaseous mixture velocity

In order to determine the intrinsic parameters of the gas-solid reaction, it is necessary to minimize the effect of external mass transfer phenomena. This can be achieved by using a high velocity of reactive gases such that the reaction rate becomes unaffected. As the mass transfer phenomena are promoted at high temperature, the effect of the gas velocity on the reaction rate of a chlorination reaction was studied at high temperature \cite{2} or at two different temperatures. The experimental results of the effect of Cl\(_2\)+O\(_2\) velocity on the oxychlorination of Fe\(_2\)O\(_3\) at 750 \, ^\circ\text{C} and 950 \, ^\circ\text{C} are given in figure 6. They indicate that the gas velocities

\[\text{Figure 5. Evolution of \% weight change as function of time during isothermal treatment of MgO in Cl}_2+\text{O}_2 \text{ at various chlorine contents.}\]

\[\text{Figura 5. Evolución del \% de cambio de peso como función del tiempo durante el tratamiento isotérmico de MgO en atmósfera de Cl}_2+\text{O}_2 \text{ con diferentes contenidos de cloro.}\]

\[\text{Figure 6. Oxychlorination rate of Fe}_2\text{O}_3 \text{ versus gas velocity of Cl}_2+\text{O}_2 \text{ at (a) 750 \, ^\circ\text{C} and (b) 950 \, ^\circ\text{C}.}\]\n
\[\text{Figura 6. Tasa de la volatilización por oxicloraación del Fe}_2\text{O}_3 \text{ en relación a la velocidad de un fluido gaseoso de Cl}_2+\text{O}_2 \text{ (a) 750 \, ^\circ\text{C y (b) 950 \, ^\circ\text{C}.}\]
higher than 0.6 cm/second are sufficient to minimize the effect of external mass transfer at both chosen temperatures. Consequently, all kinetic parameters of hematite oxychlorination were investigated by using a gases velocity of about 0.83 cm/s.

4.4. Effect of the reactive gases composition on the reaction rate

The gaseous mixtures used in the chlorination studies were Cl$_2$+CO, Cl$_2$, and Cl$_2$+O$_2$. As it could be expected, the rate of a chlorination reaction will be affected by the ratio of these gases in the chosen gaseous mixture.

The following examples demonstrate the effects of the Cl$_2$/CO and Cl$_2$/O$_2$ ratios on the chlorination reaction rates of several oxides. Experimental results are summarized in figure 7. The results of carbochlorination rate of Cr$_2$O$_3$ at 800 °C, as a function of Cl$_2$/(Cl$_2$+CO) molar ratio is shown in figure 7 a). The reaction rate of Cr$_2$O$_3$ with 100 % CO is almost equal to zero and rises when the chlorine content in the gas mixture increases. It reaches a maximum value for a Cl$_2$/(Cl$_2$+CO) ratio of ≈0.5. Beyond this ratio, the reaction rate decreases as the chlorine content increases and reaches a minimum.

![Figure 7](image-url)

**Figure 7.** Evolution of the reaction rates as a function of the reactive gases ratios for: (a) carbochlorination of Cr$_2$O$_3$ at 800 °C; (b) oxychlorination of Cr$_2$O$_3$ at 800 °C and Fe$_2$O$_3$ at 750 °C and (c) oxychlorination of MgO at 950 °C.

**Figura 7.** Evolución de las tasas de reacción en función de cocientes de reacción de los gases para: (a) carbocloración de Cr$_2$O$_3$ a 800 °C; (b) oxicloraclión de Cr$_2$O$_3$ a 800 °C y Fe$_2$O$_3$ a 750 °C y (c) oxicloraclión de MgO a 950 °C.
value with 100% Cl2. The observed maximum reaction rate at Cl2/(Cl2+CO) ratio of 0.5 suggests that the carbochlorination of Cr2O3 with Cl2+CO follows equation (1). The effect of the Cl2/(Cl2+O2) molar ratio on the oxychlorination of Cr2O3 at 800 °C is represented in figure 7 b). The oxychlorination rates passes by a maximum corresponding to a Cl2/(Cl2+O2) molar ratio of about 0.8. This value corresponds to a molar ratio of Cl2/O2 equal to 4. These results show that chlorination of Cr2O3 with Cl2+O2 gas mixture takes place according to equation (3) involving the formation of chromium oxychloride as the main reaction product.

\[
\frac{1}{2} \text{Cr}_2\text{O}_3 + \text{Cl}_2 + \frac{1}{4} \text{O}_2 \rightarrow \text{CrO}_2\text{Cl}_2 \quad (3)
\]

As this could be expected, the oxychlorination rate of Fe2O3 (Fig. 7 b)) increases when the Cl2/(Cl2+O2) molar ratio augments and it reaches a maximum value for the chlorination in absence of oxygen. Thus chlorination of Fe2O3 can be described by equation (4).

\[
\frac{1}{3} \text{Fe}_2\text{O}_3 + \text{Cl}_2 \rightarrow \frac{2}{3} \text{FeCl}_3 + \frac{1}{2} \text{O}_2 \quad (4)
\]

Similarly, the reaction rate of MgO with Cl2+O2 (Fig. 7 c)) augments with the decrease of oxygen content in the Cl2+O2 gas mixture. This may be explained by the fact that the presence of oxygen in the gas mixture will shift the reaction of equation (2) to the left side.

4.6. Determination of the apparent activation energy of reaction

4.6.1. Determination of \( E_a \) of the reaction from isothermal treatment

The apparent activation energy \( E_a \) could be evaluated from Arrhenius equation:

\[
\ln(\text{Reaction rate}) = \frac{E_a}{RT} - \ln(A)
\]

As indicated by figure 8, the apparent reaction orders for Cr2O3 oxychlorination at 800 °C with respect to Cl2, O2, and Cl2+O2 are respectively about 1.08, 0.23 and 1.29. One may note that the presence of oxygen enhances the reaction rate since \( n_{\text{O}_2} = 0.23 \). This observation agrees well with the oxychlorination of Cr2O3 (Eq. (3)) that consumes oxygen to generate chromium (VI) oxychloride (CrO2Cl2).

As a contrast, the apparent reaction order with respect to Cl2, O2, and Cl2+O2 for MgO oxychlorination were 0.98, –0.37 and 0.65, indicating that the oxygen had a negative effect on the chlorination rate of MgO. This result was expected and agrees with equation (2). On the other hand for both the studied cases (Cr2O3 or MgO), the global order is almost equal to the algebraic sum of the partial orders: \( n_{(\text{Cl}_2+\text{O}_2)} = n_{\text{Cl}_2} + n_{\text{O}_2} \).

4.5. The apparent reaction orders with respect to the reactive gases

4.5.1. Apparent reaction orders with respect to Cl2 and O2

To determine the apparent reaction order with respect to Cl2, a series of experimental tests were carried out at constant partial pressure of oxygen (0.33 atm) and the chlorine partial pressure was varied in a defined interval. Similarly, the apparent reaction order with respect to O2 was derived from the experimental results at constant Cl2 partial pressure (0.33 atm). In all cases, the chosen gas velocities correspond to optimal values defined previously. Nitrogen was used as a carrier gas.

4.5.2. Apparent reaction order with respect to Cl2+O2

A gas mixture of Cl2+O2+N2 with a velocity of 0.55 cm/second was used. The Cl2/O2 molar ratio of the gas mixture was kept constant at 4, while the partial pressure of Cl2+O2 was varied from 0.38 to 1.00 atm.

Figure 8. Evolution of the reaction rate of Cr2O3 oxychlorination at 800 °C versus partial pressure of reactive gases.

Figura 8. Evolución de la tasa volatilizada en la reacción de oxicloración del Cr2O3 a 800 °C frente a la presión parcial de los gases reaccionantes.
where $R$ is the gas constant, $T$ is the temperature in Kelvin, and $k$ is a constant.

To determine $E_a$, the solid is first heated at a predetermined temperature under nitrogen atmosphere, and when a steady state is reached, the gas mixture is introduced to react with the solid at constant temperature and the evolution of the reaction extent as a function of time is recorded. Therefore, the initial reaction rate for every tested temperature is determined. The value of $E_a$ is deduced from the slope of the straight line representing equation (6).

4.6.2. Geometry of interface (shape factor)

As the reaction of solid with chlorine takes place generally at interface of two phases, the geometry of this interface plays an important role to localize the reaction front and consequently affecting the reaction progress. The mathematical formulations of the reaction progress (Eqs. (7) – (13)) are given for different shapes of particles. These formulations are designed as conversion functions of solid 'g(X)'. The interpretations of these functions were published by Szekely J. et al.[11]. One may note that these formulations are valid when the overall reaction rate is affected by the slowest step.

Equation (7) is considered to describe a reaction controlled by the chemical reaction in the case of shrinking nonporous particles (with or without a solid porous product) and porous particles with unchanged overall size. It also applies for a mechanism affected by pore diffusion in the case of complete gasification of porous solids.

\[ 1-(1-X)^{1/F_p} = kt \]  
\[ X = kt \text{ for } F_p = 1 \]  
\[ 1-(1-X)^{1/2} = kt \text{ for } F_p = 2 \]  
\[ 1-(1-X)^{1/3} = kt \text{ for } F_p = 3 \]  
\[ X^2 = kt \text{ for } F_p = 1 \]  
\[ X+(1-X)\ln(1-X) = kt \text{ for } F_p = 2 \]  
\[ 1-3(1-X)^{1/3}+2(1-X) = kt \text{ for } F_p = 3 \]  

where $k$ : constant, $t$ : reaction time, $X$ : extent of reaction (ratio of weight of the reacted fraction to initial weight), $F_p$ : particle shape factor (1, 2, and 3 for infinite slabs, long cylinders, and for spheres).

Equations (11) through (13) are pertinent to pore diffusion control in reaction of porous solids or in nonporous solids with formation of a porous solid product.

The geometry of the solid particles combined with the value of the apparent activation energy permit to suggest the rate-controlling step of the chlorine-solid reactions. The following paragraphs describe several examples.

Figure 9 groups the data obtained during the isothermal carbochlorination of Cr$_2$O$_3$ between 500 and 900 °C[2]. The apparent activation energy is 100±2 kJ/mol (Fig. 9 a)). This value tends to indicate that the overall rate of Cr$_2$O$_3$ carbochlorination in Cl$_2$+CO is controlled by the chemical reaction. To check this hypothesis, the mathematical formulation of the experimental data was attempted using equations (7) through (13). The best result (Fig. 9 b) and c)) of data linearization, with a correlation coefficient of $\approx 0.998$[10], was obtained using equation (10). This supports the hypothesis that the carbochlorination reaction of Cr$_2$O$_3$ particles is controlled by a chemical reaction according to the shrinking sphere model described by equation (10).

4.7. Chlorination kinetics of combined oxides

The oxychlorination kinetics of chromite could be presented as a case study. The chromite presents a spinel structure including Cr, Fe, Mg, and Al oxides. As mentioned in section 3, the chromite sample used had the following composition: (Fe$^{2+}_{0.30,}Mg_{0.70}$) (Cr$^{3+}_{1.56},Al_{0.20,}Fe^{3+}_{0.07})O_4$. This solid can also be formulated as: 30.9 % FeCr$_2$O$_4$, 51.0 % MgCr$_2$O$_4$, 13.7 % MgAl$_2$O$_4$, and 4.4 % Fe$_3$O$_4$. The temperature effect on the oxychlorination of chromite was explored between 600 and 1,050 °C[2 y 6]. Results showed that the iron associated to chromium reacted with Cl$_2$+O$_2$ starting at 600 °C. This reaction could be considered as that of the simple spinel FeCr$_2$O$_4$, while MgCr$_2$O$_4$ reacts at temperature higher than 900 °C and MgAl$_2$O$_4$ seems to be refractory to the oxychlorinating gas mixture up to 1,050 °C. Figure 10 a) represents the evolution of % weight change versus time of the chromite sample during its reaction with Cl$_2$+O$_2$ between 925 and 1,050 °C. This figure
shows the sharp change of the reaction rate of chromite at X ≈ 0.35. The first part of the weight loss corresponds essentially to the chlorination of FeCr$_2$O$_4$ + Fe$_3$O$_4$ (LCh) while the second part represents the reaction of MgCr$_2$O$_4$ with Cl$_2$+O$_2$.

The temperature coefficient of the reaction was determined using the Arrhenius formula. It was interesting to follow the evolution of apparent activation energy as a function of reaction extent. The variation of $E_a$ values versus the reaction extent is shown in figure 10 b) for the temperatures range of 925 to 1050 °C.

Figure 10 b) shows that the value of $E_a$ for X comprising between 0.05 and 0.30, is lower than 60 kJ/mol. This is followed by a sharp increase of the apparent activation energy to 260 kJ/mol for X > 0.3.

To get an insight of the reactions of chromite with Cl$_2$+O$_2$, a detailed study of the effects of temperature and partial pressures of gases [$n$(Cl$_2$+O$_2$) = $n$Cl$_2$+$n$O$_2$] on the oxychlorination of chromite and its simple oxides (Cr$_2$O$_3$, Fe$_2$O$_3$, and MgO) was performed. Results are summarized in figure 11 and in table I. Although simple oxides are included in the spinel structure of chromite, the following hypothesis can be made:

- to chlorinate the chromite constituents (FeO,Cr$_2$O$_3$ and/or MgO,Cr$_2$O$_3$) their individual oxides have to react with the chlorinating gas mixture,
- the global chlorination rate of chromite constituents depends of the slowest reaction rate of the individual oxide,

**Figure 9.** Plots of data obtained during isothermal carbochlorination of Cr$_2$O$_3$ in Cl$_2$+CO: (a) Arrhenius diagram of carbochlorination and (b) and (c) Mathematical fitting of the experimental data using equation (10).

**Figura 9.** Gráficos de los datos obtenidos durante la carbocloración isoterma del Cr$_2$O$_3$ en Cl$_2$+CO: (a) Diagrama de Arrhenius de carbocloración y (b) y (c) Ajustes matemáticos de los datos experimentales usando la ecuación (10).
the kinetics parameters of the chlorination of chromite constituents should have comparable values to those of chlorination of individual oxide having the slowest reaction rate.

Figure 11 shows that the oxychlorination of chromite for \( X \leq 0.25 \) (FeCr\(_2\)O\(_4\)) is characterized by a value of \( E_a \approx 151 \text{ kJ/mol} \) between 600 and 825 °C. A similar value is obtained for the oxychlorination of Fe\(_2\)O\(_3\).

Figure 10. Oxychlorination of chromite between 925 and 1050 °C in Cl\(_2\)+O\(_2\) atmosphere. (b) Shows the evolution of the apparent activation energy values as a function of the reaction extent calculated from the data of (a).

Figura 10. Evolución del % de cambio de peso frente al tiempo de la oxiclaronación de la cromita por atmósferas de Cl\(_2\)+O\(_2\). La figura (b) indica la evolución de los valores de la energía aparente de activación en función de la extensión de la reacción que se puede calcular a partir de los datos expresados en la figura (a).

Figure 11. Arrhenius diagrams for the oxychlorination of chromite and its simple oxides in Cl\(_2\)+O\(_2\).

Figura 11.- Diagramas de Arrhenius de la oxiclaronación de cromita y sus óxidos sencillos asociados en atmósfera de Cl\(_2\)+O\(_2\).
Moreover, the oxychlorination rates of \( \text{FeCr}_2\text{O}_4 \) and \( \text{Fe}_2\text{O}_3 \) are identical. In addition, the apparent reaction orders with respect to \( \text{Cl}_2+\text{O}_2 \), \( \text{Cl}_2 \), and \( \text{O}_2 \) for the oxychlorination of chromite and \( \text{Fe}_2\text{O}_3 \) have the same tendency (Table I). While the kinetics parameters of \( \text{Cr}_2\text{O}_3 \) oxychlorination are different from those obtained for the chromite (Fig. 11 & Table I).

These experimental results indicate that the oxychlorination rate of iron oxides governs the global rate of the \( \text{FeCr}_2\text{O}_4 \) reaction with \( \text{Cl}_2+\text{O}_2 \) for temperatures lower than 825 °C.

At temperature higher than 825 °C, the oxychlorination of chromite (\( X \leq 0.25 \)) proceeds with a value of \( E_a \approx 57 \text{ kJ/mol} \). This value and the reaction rate are fairly similar to those obtained for \( \text{Cr}_2\text{O}_3 \) oxychlorination (\( E_a \approx 46 \text{ kJ/mol} \)). The positive value of \( n_{\text{O}_2} \) for a determined reaction extent (Table I) reinforces the hypothesis that the oxychlorination rate of chromite (\( X \leq 0.30 \)), at \( T > 825 \) °C is controlled, fully or partly, by the oxychlorination rate of \( \text{Cr}_2\text{O}_3 \).

The temperature coefficient of the chromite chlorination for \( X > 0.4 \) (\( \text{MgO}, \text{Cr}_2\text{O}_3 \)) is about 262 kJ/mol from 925 to 1050 °C. On the other hand, the values of \( E_a \) for \( \text{Cr}_2\text{O}_3 \) (\( E_a = 46 \text{ kJ/mol} \)) and \( \text{MgO} \) (\( E_a = 214 \text{ kJ/mol} \)) oxychlorination as well as those of the apparent reaction orders (Table I) suggest that the oxychlorination rate of MgO controls the global rate of chromite oxychlorination for \( X \geq 0.4 \) at 925-1,050 °C.

This study shows that the oxychlorination of chromite proceeds in two stages characterized by different reactivities of chromite towards \( \text{Cl}_2+\text{O}_2 \) gaseous mixture. The first stage is the \( \text{FeCr}_2\text{O}_4 \) chlorination and the second one is that of \( \text{MgCr}_2\text{O}_4 \) chlorination. In both cases, the global reaction rate was affected by the slowest rate of oxychlorination of individual oxides (\( \text{Fe}_2\text{O}_3, \text{Cr}_2\text{O}_3, \text{MgO} \)). However, one should take into account the bond energy of simple constituents of chromite to confirm the above mentioned mechanism.

### 4.8. Comparison of the \( E_a \) derived from isothermal and non-isothermal conditions

The following paragraphs describe the theoretical approach[12] for the determination of the activation energy of a gas-solid reaction from a single TGA using linear temperature rise (LTR).

The reaction rate of a gas-solid reaction could be expressed according to equation (14):

\[
dX/dt = k_f(p_A)f_2(X)f_3(d)
\]

where \( X \) is the reaction extent, \( p_A \) is the partial pressure of the reactive gas(es) and \( d \) is the mean particle size.

The effects of partial pressure of the reactive gases, reaction extent and particle size on the reaction rate are described by \( f_1(p_A), f_2(X) \) and \( f_3(d) \), respectively.

The temperature effect on the reaction rate is given by Arrhenius law (equation (15), using the rate constant \( k \)):

\[
k = k_0 \exp \left( -E_a/RT \right)
\]
Integration of Eq. 14 in isothermal conditions gives:

\[
\int_{X_0}^{X} dX/f_2(X) \equiv g(X) = \int_{0}^{t} kf_1(p_A)f_3(d)dt
\]  

(16)

The conversion function of the solid \( g(X) \) depends generally on the geometric change occurring in the solid as the reaction proceeds. The different types of \( g(X) \) were summarized in equations (7)-(13).

If the temperature is raised linearly with respect to time the temperature evolution may be given by equation (17):

\[
T = \Phi t
\]

or

\[
dT = \Phi dt
\]

(17)  

(18)

Substituting equations (15) and (17) in equation (14) and rearranging the resulting equation, it was obtained:

\[
\ln[(dX/dT)g'(X)] = \ln[k_0f_1(p_A)f_3(d)/\Phi] - \frac{E_a}{RT}
\]

(19)

In the above equation, \( g'(X) \) is the derivative of \( g(X) \) with respect to \( X \). The activation energy can be determined from the slope of data fitted by equation (19) when the appropriate type of \( g(X) \) is determined.

4.8.1. Example of the \( E_a \) evaluation from one TGA test using LTR.

The determination of the temperature effect on the reaction of MgO with the gas mixture of \( \text{Cl}_2 + \text{O}_2 \) (\( \text{Cl}_2/\text{O}_2 = 4 \)) is chosen as example. As the reactivity of MgO towards oxychlorinating gas mixture was relatively low [2, 9 y 10] the heating rate of the sample was about 1 °C/min. Figure 12 represents the obtained results concerning the oxychlorination of MgO from 575 to 1,025 °C. Magnesium oxide started to react with the oxychlorinating gas mixture at temperatures higher than 800 °C and about 70 % of the sample had reacted at 1,025 °C.

The oxychlorination reaction of MgO with \( \text{Cl}_2+\text{O}_2 \) (\( \text{Cl}_2/\text{O}_2 = 4 \)) was also studied in isothermal conditions between 850 and 1,025 °C [8-10] in order to compare the results with those obtained using LTR method. Figure 13 traces the evolution of % weight change of the sample as a function of time during isothermal oxychlorination of MgO. The evolution of the reaction extent (\( X \leq 0.7 \)) as function of time for all isotherms was appropriately described by equation (9) [8 y 19]. Figure 14 a) represents the Arrhenius diagram giving the evolution of the natural logarithm of the rate constant “k” as a function of the inverse of temperature. A value of \( E_a \) about 213 kJ/mol was deduced from the isothermal treatment of MgO in \( \text{Cl}_2+\text{O}_2 \).

Using equation (19), it is clear that the activation energy could be determined only after finding the appropriate form of \( g(X) \). As for the isothermal tests, different models of the conversion function \( g(X) \) were tried and the best non-isothermal data fitting was obtained by using equation (9). The data were plotted in figure 14 b) using equation (19) for \( X \leq 0.70 \). A value of \( E_a \approx 215 \) kJ/mol was deduced for the non-isothermal oxychlorination of MgO between 850 °C and 1,025 °C. Figure 14 indicates that the activation energy values calculated using the two sets of data are almost identical. These results confirm the utility of non-isothermal measurements using LTR to determine the activation energy of gas-solid reactions.

5. CONCLUSIONS

The following general conclusions can be drawn from the kinetics studies of chlorination reactions of different materials through thermo-gravimetric measurements.

— The reactivity of different solids towards chlorinating gases varied widely e.g. the reaction \( \text{Cl}_2 \) with \( \text{CuFeS}_2 \) started at room
The effect of external mass and heat transfer on the reaction rate could be minimized by choosing adequate gas velocities.

The apparent reaction orders with respect to reactive gases (Cl₂, O₂, CO), obtained experimentally, were fractions, either positive or negative numbers. Although they differ from stoichiometric coefficients of the reaction, their positive or negative value depends on the considered chemical reactions. The apparent reaction order with respect to O₂ is 0.23 for Cr₂O₃ oxychlorination, while that of MgO oxychlorination nO₂ ≈ –0.37. The oxychlorination reactions of MgO and Cr₂O₃ required a temperature ≥ 500 °C.
with Cl2+O2 explain this difference concerning
the oxygen role in the two reactions.
— The value of $E_a$ depends on the type of solid,
the nature of the chlorinating gas mixture
(Cl2, Cl2+O2, Cl2+CO), the studied
temperature range, the physical state of the
reaction products, etc.
— The isothermal and the linear temperature
raise techniques gave essentially similar kinetic
parameters. Further, from the formulation of
the reaction interface geometry, the rate
controlling step could be predicted accurately.

This study thus demonstrated that, simple thermo-
gravimetric measurements can be utilized for
measurement of complex reaction kinetics involving
gas – solid reactions such as chlorination of variety
of solid samples with number of reactive gases
compositions. The obtained kinetics information
can be used for process optimization involving
beneficiation of lean ores and extraction of several
valuable metals from their oxide or sulfide ores, which
are otherwise difficult by traditional chemical
beneficiation and/or extraction methods.

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