# Model for Computational Analysis of the Concentration of Sulphur Removed during Temperature Dependent Oxidation of Iron Oxide Ore

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**Abstract**: Model for computational analysis of the concentration of sulphur removed during temperature dependent oxidation (by potassium chlorate) of iron oxide ore has been derived. The model;

$$\%S = \left( \begin{array}{c} 0.1891 \\ \hline LogT \end{array} \right)$$

was found to predict the concentration of sulphur removed, very close to the corresponding %S values obtained from the actual experimental process. It was found that the model is dependent on the values of the treatment temperature used during the desulphurization process. The validity of the model is believed to be rooted in the expression  $[(T)^{\%S}] = /k_n$  where both sides of the expression are approximately equal to 2. The positive or negative deviation of each of the model-predicted values of %S from those of the corresponding experimental values was found to be less than 21% which is quite within the range of acceptable deviation limit of experimental results, hence showing the validity and usefulness of the model for computational and predictive analysis.[Researcher. 2010;2(2):25-29]. (ISSN: 1553-9865)

Keywords: Model, Sulphur Removed, Iron Oxide Ore, Oxidation, Potassium Chlorate.

#### 1. Introduction

Studies (Uwadiele, 1984) revealed that Agbaja iron ore deposit is the largest known Nigerian iron ore deposit estimated at 1250 metric tonnes of ore reserve. It consists of oolitic and pisolitic structures rich in iron oxides, in a matrix that is predominantly clay. The principal constituent mineral is goethite, with minor hematite, maghemite, siderite, quartz, kaolinite pyrite and an average of 0.09%S.

Intensive and selective oil agglomeration of Agbaja iron ore has been carried out by Uwadiele (1990). The researcher, starting from the crude ore Fe content (45.6%), concentrated the ore by oil agglomeration technique to 90% Fe recovery and 65% Fe assay. He stated that the ore require grinding to minus 5 $\mu$ m to effect adequate liberation. These results were obtained at optimum pH 9. Uwadiele and Whewell (1988) studied the effect of temperature on magnetizing reduction of Agbaja iron ore. The results of the investigation showed that the fine-grained oolitic Agbaja iron ore, which is not responsive to conventional processing techniques, can be upgraded by the magnetizing reduction method with an Fe recovery of 87.3% and Fe assay of 60% at 600<sup>o</sup>C.

Attempt has been made by Kulkarni and Somasundaran (1980) to enhance concentrate Fe recovery. The researchers stated that concentrate Fe recovery decreases progressively below pH 8. In this pH region, oleate used is present as dispersion of oleic acid, and its adsorption on the surface of the iron oxides is similar to the process of hetero-coagulation involving positively charged iron oxide particles and negatively charged oleic acid droplet.

Studies by Nwoye (2008) reveal that Agbaja oolitic iron ore, which has not been responsive to so many upgrading processes, has been upgraded to 73.4% Fe assay (starting from as-received concentrate assaying 56.2%Fe) by a process referred to as pyrometallurgical-oxidation method. The researcher investigated mainly the effects of treatment temperature and oxidant (KClO<sub>3</sub>) on the upgrading process. It was established that 800°C is the optimum temperature for the upgrading step considering the range of temperature used (500-800°C). It was observed from results of the investigation that both oxidant and temperature increase (up to 12g per 50g of iron ore and maximum of 800°C respectively) during the process are vital conditions for improving on the grade of the ore concentrate.

A model for computational analysis of the concentration of iron upgraded during dry beneficiation of iron oxide ore has been derived by Nwoye et al., (2009). The model;

%Fe = 
$$2.25[(\ln (T/\mu))^{2.58}]$$
 (1)

shows that the concentration of upgraded iron is dependent on the treatment temperature T, used when the mass of iron oxide ore  $\mu$ , added is constant.

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Nwoye (2008) studied the desulphurization of Agbaja iron oxide ore concentrate by using solid potassium trioxochlorate (V) (KClO<sub>3</sub>) as oxidant and a treatment temperature range:  $500 - 800^{\circ}$ C. The results of the investigation revealed that simultaneous increase in both the percentage of the oxidant added (up to 15g per 50g of ore) and treatment temperature (maximum  $800^{\circ}$ C) used gives the ideal conditions for increased desulphurization efficiency. This translates into high desulphurization (up to 15g per 50g of ore) and treatment temperature (maximum s00°C) are high.

Further investigation (Nwoye, 2009) was on the mechanism and process analysis of desulphurization of Agbaja iron ore concentrate using powdered potassium trioxochlorate (v) (KClO<sub>3</sub>) as oxidant. Concentrates were treated at a temperature range  $500 - 800^{\circ}$ C. Results of the process analysis indicate that oxygen required for the desulphurization process was produced following decomposition of KClO3 within a temperature range 375-502°C. It was observed that this temperature range is the Gas Evolution Temperature Range (GETR) for sulphur present in Agbaja iron ore. Sulphur vapour and oxygen gas produced at this temperature range were believed to have reacted to form and liberate SO<sub>2</sub>. The process analysis suggests that the mechanism of the desulphurization process involves gaseous state interaction between oxygen and sulphur through molecular combination. The results for the extent of desulphurization reveal that simultaneous increase in both the percentage of the oxidant added and treatment temperature used (up to 15g KClO3 per 50g of ore and maximum of 800°C respectively) are the ideal conditions for the best desulphurization efficiency.

Report of the investigation by Bardenheuer and Geller (1934) indicated that the sulphur transfer from metal to slag or slag to gas during desulphurization involves oxygen transfer in the opposite direction. They posited that the mechanism of such desulphurization involves oxidation of sulphur resident in the metal or slag by oxygen from the slag through ionic exchange between the oxygen and sulphur, since the whole system is made up of liquid/molten condition during this process. They maintained that oxygen in the slag comes from CaO, which is one of the products of decomposition of CaCO<sub>3</sub> deposited into the slag as a slag forming agent.

St Pierre and Chipman (1956) studied gas-slag system during iron making. Results of their investigation show that at oxygen partial pressure below about  $10^{-5}$  atm., sulphur dissolves in the melt as sulphide ions; at oxygen partial pressure higher than  $10^{-3}$  atm., sulphur enters the melt as sulphate ions. In both cases, they stated that both the sulphide and sulphate ions leave the furnace through the slag. They therefore concluded that the mechanism of such desulphurization process is oxidation of sulphur by oxygen from the slag through ionic exchange between the two participating elements.

Turkdogan and Darken (1961) found that at a temperature well below about  $1600^{\circ}$ C, the pyrosulphate reaction also occurs. They found that this reaction was an enhancement to the desulphurization process actually taking place in the furnace. Also oxygen for this process was found to come from the slag, engaging sulphur in ionic exchange; being the mechanism of such process.

It was discovered that one of the most important factors influencing the desulphurization process during iron making is the state of oxidation of the bath (Pehlke et.al 1975).

Nwoye et al., 2009 derived a model for the predictive analysis of the concentration of sulphur removed as result of the molecular-oxygen-induced desulphurization of iron oxide ore (potassium chlorate being the oxidant). The model;

$$\%S = \left(\frac{0.0415}{\text{Log}}\right)$$
(2)

was found to predict the concentration of sulphur removed, very close to the corresponding %S values obtained from the actual experimental process. It was found that the model is dependent on the values of the weight-input of the oxidant , (KClO<sub>3</sub>) during the desulphurization process. The validity of the model is believed to be rooted in the expression  $k_n[(\ )^{\mu\%S}]$  =T/ where both sides of the expression are correspondingly almost equal. The positive or negative deviation of each of the model-predicted values of %S from those of the corresponding experimental values was found to be less than 33% which is quite within the range of acceptable deviation limit of experimental results.

A model for computational analysis of the concentration of sulphur removed during oxidation of iron oxide ore by powdered potassium chlorate has been derived by Nwoye et al., (2009). The model;

$$\%S = \left(\frac{0.0357}{\text{Log}}\right) \tag{3}$$

indicates that the predicted %S is dependent on the weight-input of KClO<sub>3</sub>, added during the desulphurization process. The maximum deviation of the model-predicted values of %S from those of the corresponding experimental values was found to be less than 37%

Nwoye et al., (2009) derived a model for predicting the concentration of sulphur removed during gaseous desulphurization of iron oxide ore . The model;

$$\%S = \left(\frac{0.0745}{\text{LogT}}\right) \tag{4}$$

shows that the predicted %S is dependent on the treatment temperature T, used during the desulphurization process.

The aim of this work is to derive a model for computational analysis of the concentration of sulphur removed during temperature dependent oxidation of Agbaja (Nigerian) iron oxide ore

#### 2. Model

The solid phase (ore) is assumed to be stationary, contains some unreduced iron remaining in the ore. It was found (Nwoye, 2008) that oxygen gas from the decomposition of KClO<sub>3</sub> attacked the ore in a gas-solid reaction, hence removing (through oxidation) the sulphur present in the ore in the form of SO<sub>2</sub>.Equations (5) and (6) show this.

$$2\text{KClO}_{3 (s)} \longrightarrow 2\text{KCl}_{(s)} + 3O_{2 (g)}$$
(5)  
$$S_{(s)}\text{Heat} S_{(g)} + O_{2 (g)} \longrightarrow SO_{2 (g)}$$
(6)

#### 2.1 Model Formulation

Experimental data obtained from research work (Nwoye,2007) carried out at SynchroWell Research Laboratory, Enugu were used for this work. Results of the experiment as presented in report (Nwoye, 2007) and used for the model formulation are as shown in Table 1.

Computational analysis of the experimental data shown in Table 1, gave rise to Table 2 which indicate that;

$$[(T)^{\%S}] = /k_n \text{ (approximately)}$$
(7)

$$k_n [(T)^{\%S}] = (8)$$

Taking logarithim of both sides

$$Log (k_n[(T) ^{\% S}]) = Log$$
(9)

$$Logk_n + Log [(T)^{\%S}]) = Log$$
(10)

$$Logk_n + \%SLogT = Log$$
 (11)

$$\% \operatorname{Slog} T = \operatorname{Log} - \operatorname{Log} k_n \tag{12}$$

$$%S = Log - Logk_n$$
(13)  
$$LogT$$

Introducing the values of ,  $k_n$  and into equation (13) (since the are constants) and evaluating further, reduces it to;

$$S = 0.1891$$
 (14)  
LogT

Therefore  

$$%S = \underline{D_e}$$
 (15)  
LogT

Where

- %S = Concentration of sulphur removed during the pyrometallurgical-oxidation process.
  - $k_n = 9.71$  (Decomposition coefficient of

KClO<sub>3</sub> relative to its weight input (15g per 50g of the iron ore) determined in the experiment (Nwoye,2007)

- ( )= 0.9989 (Temperature coefficient relative to weight-input of KClO<sub>3</sub>) determined in the experiment (Nwoye,2007)
- ( ) = Weight of KClO<sub>3</sub> added as oxidant (g)
   T = Treatment temperature used for the process (<sup>0</sup>C)
- $D_e = 0.1891$  (Assumed desulphurization enhancement factor)

Table 1: Variation of concentration of sulphur removed with treatment temperature (Nwoye,2007)

$T (^{0}C)$		%S
500	50	0.070
550	50	0.073
600	50	0.080
650	50	0.080
700	50	0.080
750	50	0.083
		1

Table

2:

Variation of  $/k_n$  with T  $^{\% S}$ 

/k <sub>n</sub>	T <sup>%S</sup>
1.5448	1.5440
1.5448	1.5841
1.5448	1.6671
1.5448	1.6778
1.5448	1.6878
1.5448	1.7312

## 3. Boundary and Initial Condition

Consider iron ore (in a furnace) mixed with potassium chlorate (oxidant). The furnace atmosphere is not contaminated i.e (free of unwanted gases and dusts). Initially, atmospheric levels of oxygen are assumed just before the decomposition of KClO<sub>3</sub> (due to air in the furnace).Weight, M of iron oxide ore used; (50g), and treatment time; 360secs. were used. Treatment temperature range;  $500-750^{\circ}\text{C}$ , ore grain size;  $150\mu\text{m}$ , and weight of KClO<sub>3</sub> (oxidant); 15g were also used. These and other process conditions are as stated in the experimental technique (Nwoye; 2007).

The boundary conditions are: furnace oxygen atmosphere due to decomposition of  $KCIO_3$  (since the furnace was air-tight closed) at the top and bottom of the ore particles interacting with the gas phase. At the bottom of the particles, a zero gradient for the gas scalar are assumed and also for the gas phase at the top of the particles. The reduced iron is stationary. The sides of the particles are taken to be symmetries.

#### 4. Model Validation

The formulated model was validated by direct analysis and comparison of %S values predicted by the model and those obtained from the experiment for equality or near equality.

Analysis and comparison between these %S values reveal deviations of model-predicted %S values from those of the experiment. This is attributed to the fact that the surface properties of the ore and the physiochemical interactions between the ore and the oxidant (under the influence of the treatment temperature) which were found to have played vital roles during the oxidation process (Nwoye, 2007) were not considered during the model formulation. This necessitated the introduction of correction factor; to bring the model-predicted %S values to those of the experimental %S values (Table 3).

Deviation (Dv) (%) of model-predicted %S values from experimental %S values is given by

$$Dv = \left(\frac{Sp - Se}{Se}\right) x \ 100 \tag{16}$$
  
Where

Sp = Predicted %S values from model Se = Experimental %S values

Correction factor (Cf ) is the negative of the deviation i.e  $% \left( {{{\bf{C}}_{{\bf{F}}}}_{{\bf{F}}}} \right)$ 

$$Cf = -Dv$$
 (17)

Therefore

$$Cf = -\left(\begin{array}{c} \underline{Sp - Se} \\ \underline{Se} \end{array}\right) \times 100$$
(18)

Introduction of the corresponding values of Cf from equation (18) into the model gives exactly the corresponding experimental %S values (Nwoye, 2007).

#### 5. Results and Discussion

The derived model is equation (14) or (15). A comparison of the values of %S from the experiment and those from the model shows minimum positive and negative deviations less than 21% which is quite within the acceptable deviation limit of experimental results hence depicting the reliability and validity of the model. This is shown in Table 3.

The validity of the model is believed to be rooted in expression  $[(T)^{\%S}] = /k_n$  where both sides of the equation are approximately equal to 2. Table 2 also agrees with equation (7) following the values  $/k_n$  and T  $^{\%S}$  evaluated from Table 1 as a result of corresponding computational analysis.

The value 0.1891 has a direct relationship with the value of %S as shown in equation (14). This indicates that the constant contributes directly (as a multiplying factor) to the predicted concentration of sulphur removed from the ore. Based on the foregoing, the constant is denoted as desulphurization enhancement factor  $D_e$ . Table 3: Comparison between %S removed as predicted by model and as obtained from experiment (Nwoye ,2007).

%S	e	%S <sub>M</sub>	Dv (%)	Cf (%)
0.	070	0.0701	+0.14	-0.14
0.	073	0.0690	-5.48	+5.48
0.	080	0.0681	-14.88	+14.88
0.	080	0.0672	-16.00	+16.00
0.	080	0.0665	-16.88	+16.88
0.	083	0.0658	-20.72	+20.72

Where

 $S_e = S$  values from experiment (Nwoye,2007)  $S_p = S$  values predicted by model

## 6. Conclusion

The model computes the concentration of sulphur removed during temperature dependent oxidation of Agbaja iron oxide ore using powdered potassium chlorate. The validity of the model is believed to be rooted in the expression  $[(T)^{\% S}] = /k_n$  where both sides of the equation are approximately equal to 2. The deviation of the model-predicted %S values from those of the experiment is less than 21% which is quite within the acceptable deviation limit of experimental results.

Further works should incorporate more process parameters into the model with the aim of reducing the deviations of the model-predicted %S values from those of the experiment

#### Acknowledgement

The authors thank Dr. Ekeme Udoh, a modelling expert at Linkwell Modelling Centre Calabar for his technical inputs. The management of SynchroWell Nig. Ltd. Enugu is also appreciated for permitting and providing the experimental data used in this work.

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## 12/9/2009