ESTUDO DA FORMAÇÃO E REDUÇÃO DA FERRITA DE ZINCO CONTIDAS NOS PÓS DE ACIARIA ELÉTRICA USANDO MISTURAS CO-CO₂*

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Resumo
O presente trabalho trata da formação da ferrita de zinco a partir da mistura e posterior reação solido-sólido entre o óxido de zinco e óxido de ferro - Fe₂O₃ puros em proporções equimolares. A faixa de temperatura onde teve lugar a formação foi entre 1073 e 1373 K. Testes de redução da ferrita de zinco sintética-ZF e pó de aciaria elétrica-EAFD foram realizadas usando misturas de CO e CO₂ que variaram entre 50 e 100% CO. Ambos os testes (formação e redução de ZF e EAFD) foram suportados usando técnicas de caracterização física, química, estrutural e microscópica. No caso da redução, foi observado que tanto a temperatura como a percentagem de CO, foram os fatores que, mais afetaram a redução de ferrita de zinco contida no EAFD. A máxima percentagem de redução foi 85%, para 100% CO, 1373 K e 105 min, e 52%, para 50% CO, 1373 K e 105 min. Por outro lado, atingiu o 100% para 100% CO, 1273K, em 80.75 min, 60%, para 50% CO a 1373 K durante 105 min.

Palavras-chave: Redução; Pós de aciaria elétrica; Ferrita de zinco.

STUDY OF FORMATION AND REDUCTION OF ZINC FERRITE CONTAINED ELECTRIC ARC FURNACE DUSTS BY CO - CO₂ GAS MIXTURES

Abstract
The present work deals about the zinc ferrite synthesis, occurred through a solid-solid reaction in a selected range of temperatures, using as reactant an equimolar mixture of pure iron oxide- Fe₂O₃ and pure zinc oxide - ZnO. The range temperature of formation was of 1073 to 1373 K. After that, took place the reduction of equimolar synthetic zinc ferrite-ZF and Electric Arc Furnace Dusts-EAFD by gas mixtures CO - CO₂, from 50% and 100% of CO. Both testes (formation and reduction) were supported by physical, chemical, structural and microscopic characterizations of both, initial zinc ferrite generated in laboratory and the remained after reaction. It was observed that the temperature and CO content were the main factors affecting the zinc ferrite reduction. The maximum reductions of ZF indexes obtained in these experiments were 85%, for 100% CO at 1373 K, in 105 min, and 52%, for 50% CO at 1373 K, elapsed 105 min and 100%, for 100% CO at 1273 K, in 80.75 min, and 60%, for 50% CO at 1373K during 105 minutes.

Keywords: Reduction; EAFD; Zinc ferrite.

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* Contribuição técnica ao 45º Seminário de Redução de Minério de Ferro e Matérias-primas, 16º Simpósio Brasileiro de Minério de Ferro e 3º Simpósio Brasileiro de Aglomeração de Minério de Ferro, parte integrante da ABM Week, realizada de 17 a 21 de agosto de 2015, Rio de Janeiro, RJ, Brasil.
1 INTRODUCTION

Several residues generated in the reduction and refining sectors of a steelwork plant, bears expressive quantities of zinc in their compositions, mainly in the form of oxide and ferrite compounds. Nowadays, these particulates, regarded as ambientally deleterious materials, are being the main focus of several stocking procedures and studies to reclaim its zinc content. Further, the zinc content in these residues could be as high as 25% to 40% of their composition, being commonly distributed as 70% in the form of zinc ferrite –ZF and the rest as zinc oxide and metallic one (minor part) [1]. Moreover, its quantity increases the particulates content in the generated fumes, as the recycling of galvanized steel scrap increases as burden in the steelmaking processes. Therefore, the presence of zinc ferrite in steelmaking dusts is a hard reality, and, on top of that the reduction of this substance is still not well understood. According to Jyh-Jen [2] et al., in the temperature range of 1073 to 1473K the carbothermic reduction of zinc ferrite has an apparent activation energy of circa 92.91kJ/mol. They also proposed a reduction phenomenology involving basically two steps: firstly, the initial decomposition of zinc ferrite, ZnFe₂O₄, to ZnO and Fe₂O₃, and secondly, a simultaneous and competitive final reduction of these oxides. Additionally, depending on the reduction temperature, their proposal is that the metallic zinc released should evolve as vapor, generating a degree of porosity in the solid which increases the specific surface area of the remained reacting material and consequently, its volume. This increase in volume, accompanying the reduction of the iron oxides in the final stage of the process, should also be followed by the iron sinterization, thus producing a decreasing of the pores diameter and the particles volume.

This work is a part of a broader project which major purpose is to establish a complete kinetic and morphological phenomenology, aiming a better understanding of the reduction process of the zinc ferrite present in the steelmaking dusts by gaseous mixtures of CO and CO₂.

2 METHODOLOGY

Initially, synthetic zinc ferrite samples were made in laboratory from Fe₂O₃ / ZnO: 1 / 1 mixtures following the technique developed by Hsi-Kuei Chen; Ching-Yi Yang [1] Bid S.; Pradhan S.K. [3] and Gómez-Marroquín M.C.; D'Abreu J.C. [4-9]. In continuing it was agglomerated into the shape of cylindrical briquettes according to Hui-Ning Zhang [10].

Firstly, the zinc ferrite and the Electric Arc Furnace Dusts-EAFD were structurally characterized through X-Ray diffraction, using a diffractometer Siemens, model D5000 and the following operational conditions: I=30 A, V=40 kV and copper anode (λ=1,5406 Å, CuKα). The analytical software applied was the Diffract Plus, Topas, version 2.1 from Bruker AXS for quantitative analysis by the Rietveld method. The samples of zinc ferrite and the Electric Arc Furnace Dusts-EAFD were also characterized morphologically using a Scan Electron Microscope – SEM (DSM 960 Zeiss West Germany, V = 20 kV) and a Transmission Electron Microscope – TEM (Model JEOL 2010 V = 200 kV).

The physical characterization of the zinc ferrite was made employing:

* A Multipycnometer quanta chrome V=120V and P=18psi, for the density measurements;
A Mastersize μ, Version 2.12 from Malvern Instruments, for the average size distribution and specific surface area determination. After those essays, portions of zinc ferrite and the Electric Arc Furnace Dusts-EAFD powder weighting 7 g were agglomerated in cylindrical briquettes with a diameter of 2.54 cm.

The reduction runs were performed in a tubular electric furnace, linked to a gas line with mixtures of CO, CO₂ and N₂. The experiment temperatures were 1073, 1173, 1223, 1273 and 1373K and the reaction times 8, 32.25, 56.5, 80.75 and 105 minutes. The gaseous compositions utilized were 100% CO and 50% CO – 50% CO₂.

3 RESULTS AND DISCUSSION

3.1 Structural Characterization

In Figure 1 (a) Rietveld adjusted diffractogram of a zinc ferrite synthetic sample is presented, showing the phases and their corresponding percentages: zinc ferrite (ZnFe₂O₄) – 94.82%, Iron (III) Oxide (Fe₂O₃) – 3.48%, Zinc Oxide (ZnO) – 1.70% and (b) Rietveld adjusted diffractogram of a EAFD sample is showed with: zinc ferrite (ZnFe₂O₄) – 30.82%, Iron (III) Oxide (Fe₂O₃) – 61.23%, Zinc Oxide (ZnO) – 7.95%.

3.2 Chemical Characterization

Samples of EAFD generated in a Brazilian mini-mill steelwork were chemical characterized in situ using Absorption Atomic Equipment. In Table 1 the EAFD’s chemical composition by compounds and elements are observed.
Table 1. EAFD’s chemical composition by compounds and elements

<table>
<thead>
<tr>
<th>Compound</th>
<th>SiO₂</th>
<th>MnO</th>
<th>P₂O₅</th>
<th>Cr₂O₃</th>
<th>Na₂O</th>
<th>K₂O</th>
<th>“Fe₂O₃”</th>
<th>“ZnFe₂O₄”</th>
<th>CdO</th>
<th>PbO</th>
<th>Al₂O₃</th>
<th>C</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2.81</td>
<td>1.87</td>
<td>0.28</td>
<td>0.40</td>
<td>3.77</td>
<td>3.85</td>
<td>44.04</td>
<td>1.19</td>
<td>1.98</td>
<td>28.00</td>
<td>0.06</td>
<td>2.50</td>
<td>0.57</td>
</tr>
<tr>
<td>Fe</td>
<td>44.92</td>
<td>11.79</td>
<td>1.31</td>
<td>1.45</td>
<td>0.06</td>
<td>0.14</td>
<td>1.40</td>
<td>1.60</td>
<td>0.72</td>
<td>1.42</td>
<td>0.05</td>
<td>2.32</td>
<td>0.15</td>
</tr>
</tbody>
</table>

3.3 Microscopic Characterization

Completing the identification and characterization of the phases present in the zinc ferrite samples, the SEM and TEM analysis were made. In Figure 2 (a) SEM image of the zinc ferrite, as produced in the laboratory, is presented. In it is clear the major presence of zinc ferrite white globular, besides of some pores and (b) Scanning electron microscopy (SEM) with X-ray energy dispersive spectrometry (EDS) was performed to gain further knowledge of the EAFD particles structure, morphology and their chemical composition. The same samples were also subjected to the element distribution analysis (O, Fe, Zn, Mg, Ca, Cr, Si, Mn) through X-ray mapping analysis via SEM. As for the TEM image, in Figure 3, one can observe the presence of tiny zinc ferrite particles having sizes on the range of 100 to 200 nm.

![Figure 2. SEM image of typical structures of (a) produced equimolar zinc ferrite-ZF and (b) EAFD.](image)

![Figure 3. TEM image of typical structures of produced equimolar zinc ferrite-ZF.](image)
3.4 Physical Characterization

In this aspect the samples particles of zinc ferrite-ZF and the Electric Arc Furnace Dusts-EAFD average size, specific surface area, density and porosity were measured. The results were showed in the Table 2:

<table>
<thead>
<tr>
<th>SAMPLES</th>
<th>ZF</th>
<th>EAFD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particle (clusters) average diameter, µm</td>
<td>3.95</td>
<td>1.58</td>
</tr>
<tr>
<td>Specific surface area, m²/g</td>
<td>38.23</td>
<td>54.49</td>
</tr>
<tr>
<td>Density, g/cm³</td>
<td>5.71</td>
<td>5.42</td>
</tr>
<tr>
<td>Sample’s briquette porosity</td>
<td>0.45</td>
<td>0.44</td>
</tr>
</tbody>
</table>

One must observe that the density and the sample’s briquette porosity were significantly similar to the one presented in the Tong, Lee Fui [11] work (being the theoretical zinc ferrite density is circa 5.34 g/cm³).

3.5 Reduction

The gas compositions and temperatures, elected for the tests, were chosen taking into account the phase predominance operational diagram – PPOD for the Zn-Fe-C-O system, determined by the authors [11]. In Figure 4 the PPOD for Zn-Fe-C-O is depicted. In it the equilibria ZnO-Zn and Fe₂O₃-Fe₃O₄-FeO-Fe, with a CO-CO₂ atmosphere, and the predominance area for various species, are signaled. The reduction mechanism was the reaction occurring via gaseous intermediates (equimolar zinc ferrite reduction by CO together with the Boudouard reaction):

- Indirect reductions by CO:
  3(ZnO.Fe₂O₃) (s) + CO(g) = (ZnO)₃ (Fe₃O₄)₂(s) + CO₂(g)
  (ZnO)₃(Fe₃O₄)₂(s) + 2CO(g) = (ZnO)₃(FeO)₆(s) + 2CO₂(g)
  (ZnO)₃(FeO)₆(s) + 9CO(g) = 3Zn(g) + 6Fe(s) + 9CO₂(g)

- Global indirect reduction:
  ZnO.Fe₂O₃(s) + 4CO(g) = Zn(g) + 2Fe(s) + 4CO₂(g)

- Boudouard reaction
  CO₂(g) + C(s) = 2CO(g)
The obtained kinetic curves for the zinc ferrite reduction, for different temperatures, are presented in Figure 5 (50% CO–50% CO₂ mixtures) and in Figure 6 (100% CO).

**Figure 4.** Phase predominance operational diagram – PPOD, for the Zn-Fe-C-O system.

**Figure 5.** Reduction by 50% CO - 50% CO₂ gas mixtures at different temperatures of (a) ZF and (b) EAFD.

**Figure 6.** Reduction by 100% CO gas mixtures at different temperatures of (a) ZF and (b) EAFD.
The maximum zinc ferrite synthetic reductions indexes obtained in these experiments were 85%, for 100% CO at 1373K, in 105 min, and 52%, for 50% CO at 1373K, and 100%, for 100% CO at 1373K, in 105 min, and approximately 60%, for 50% CO at 1373K elapsed 105 min, in the case of EAFD reduction. According to the information of Figure 6, it is notably that maximum reduction percentage that was achieved by EAFD´s reduction through CO pure was of circa of 100% at 1273K during a heating time of 80.75 minutes.

Another hand, Random Nucleation Reaction Model: \( kt = -\ln(1 - \alpha') \); where the \( \alpha \) is the fractional conversion, \( k \) is the specify rate constant and \( t \) is the heating time; was used in order to fit the experimental fractional conversions (reduction percentage) and so was estimated the values of Apparent Activation Energy-Ea and maximum reduction of ZF and EAFD respectively. This data is showed in Table 3.

<table>
<thead>
<tr>
<th>Samples</th>
<th>50% CO-50% CO₂</th>
<th>100% CO</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( \text{E}_a ), kJ.mol(^{-1} )</td>
<td>% Red. Max.</td>
</tr>
<tr>
<td>ZF</td>
<td>91.71</td>
<td>52</td>
</tr>
<tr>
<td>EAFD</td>
<td>100.51</td>
<td>60</td>
</tr>
</tbody>
</table>

The observed differences between the Activation Energy values, hints changing in overall reaction mechanism toward a less temperature sensitive process.

### 4 CONCLUSIONS

- The phase predominance operational diagram – PPOD, furnished the key information for temperatures and %CO values utilized in the reducing kinetic experiments;
- The microscopic characterization of the zinc ferrite powders, permitted the observation of clusters formed by the individual particles, which had the average size ranging from 100 to 200 nm;
- The physical characterization in order to determinate the density and the sample’s briquette porosity of ZF and EAFD showed values significantly similar.
- From the ZF and EAFD and gaseous CO mixtures reducing curves, the following conclusions are drawn:
  a. Increases in temperatures, reaction time and CO percentage produced, as expected, higher zinc ferrite conversions ;
  b. The Apparent Activation Energies obtained were 55.60 and 91.71 kJ/mol, for 100%CO and 50% CO-50% CO₂, and 54.74 and 100.51 kJ/mol respectively.
  c. It is notably that maximum reduction percentagem achieved by EAFD reduction through CO pure was of circa of 100% at 1273K and 80.75 minutes.
- The maximum zinc ferrite synthetic reductions indexes obtained in these experiments were 85%, for 100% CO at 1373K, in 105 min, and 52%, for 50% CO at 1373K, and 100%, for 100% CO at 1373K, in 105 min, and approximately 60%, for 50% CO at 1373K elapsed 105 min, in the case of EAFD reduction.
- To determinate a comprehensive kinetic equation for the reduction process of zinc ferrite by CO, the study under other gaseous compositions is being carried out. The results will permit a better decision making actions aiming at the reclaim of the zinc element from steelmaking dusts.
Acknowledgements

The authors wish to thank the Graduate Teaching Personnel Supporting Council – Capes and the National Research Council – CNPq, as well as Prof Guillermo S. Naranjo and Eng Raimundo N. Rodrigues Filho.

REFERENCES