Characterization and Three-dimensional Reconstruction of Pores of Self-reducing Pellets Done by EAF Dust

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This work focused on the characterization physical and microstructural of the self-reducing pellet made of Electric Arc Furnace (EAF) dust using different low cost techniques. The serial sectioning technique was used to evaluate detailed measurements of pore connections, tortuosity of pores and porosity distribution. The chemical analysis using Inductively Coupled Plasma Atomic Emission Spectroscopy (ICP-AES), Scanning Electron Microscopy (SEM), Optical Microscopy (OM) and X-ray diffraction were carried out to identify the common phases presented in EAF dust agglomerate. It was observed that the pellet phase composition is formed by iron as magnetite, metallic iron, wustite, and zinc ferrite. The visualization of the reconstructed 3D microstructure provided average qualitative and quantitative analysis of the porosity (41.61%), a consistent result and in accordance with that obtained by pycnometry technique (41.53%). As expected, these results are more precise when compared with the result obtained by two-dimensional technique (23.41%). In addition, it was calculated the value of the tortuosity parameter (0.84) that suggested a morphological structure closest to cylindrical shape.

Keywords: 3D reconstruction, serial sectioning, self-reducing pellet, EAF dust

1. Introduction

The detailed characterization of raw materials in the steel industry is recognized as an important tool to warranty the quality of steel product and largely applied on the development of new processes based on raw materials such as ultrafine iron ores, dust and solid residues from the steel industry. For instance, in agglomerates, some microstructural parameters are relevant to study the inner phenomena that take place into the reduction processes; such as the mineralogical composition, size distribution of particles, porosity and pore connectivity within these agglomerates. The raw material used to produce pellets and sinters in the iron ore sintering process can be obtained from several sources, such as, iron ore or dust recycling within the steelworks2. There are many environmental and economic benefits in recycling large amount of waste materials and energy, but, in the case of pellet and sinter produced with addition of electric arc furnace dust, it is necessary to predict the quality of the produced pellet due its impact in the production of pig iron. The chemical composition of the agglomerates will play important role in the formation of phases and consequently, in the thermodynamic and kinetic phenomena that take place into the subsequent processes. Despite of large amount of iron present in EAF dust (Fe₂O₃, FeO, Fe₂O₄, Fe particles), some impurities such as zinc, cadmium and chromium are bottleneck for the technologies that have been proposed, since high concentration of zinc leads to producing complex phases, as, zinc ferrite (ZnFe₂O₄), zinc silicates (Zn₃SiO₄) and zinc oxide (ZnO), and contains also, toxic components like lead, cadmium, arsenic and chromium, which would have environment load2. Some metallographic two-dimensional techniques like optical and electronic microscopy and X-ray diffraction have been used to characterize pellets with limited results. Wagner et al.3, developed a method to quantify porosity in iron ore pellets using digital images obtained by optical microscope. Iglesias4, characterized the sinter by digital images in optical microscope, identifying main phases like magnetite, hematite, ferrites, silicates and pores. However, in the mentioned studies and others1,2, were not considered the inherent three-dimensional characteristics of the pore structures. Rodrigues et al.5, performed a study about 3D reconstruction of the porous structure of Ti-6Al-4V alloy by serial sectioning, and the visualization of the microstructure provided a qualitative and quantitative analysis of the porosity of the alloy at different temperatures and demonstrated the feasibility of this technique to obtain detailed information regarding pore connectivity and morphology. On the other hand, the high temperature process used to treat the agglomerates of waste materials strongly depends on the diffusion of fluids within particles which is strongly affected by the porosity distributions and topology. An important parameter to estimate the porous topology is the tortuosity that reflects the material anisotropy6. The tortuosity parameter can be regarded as the relation between the effective and the linear lengths of the pore penetrated by the fluids within the reacting agglomerates. The pore diameter distributions are usually variable and can be connected and cause disorder in the fluid flow molecules inside the pore walls of the pellets. This contact generates a frictional force.

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and causes an increasing of the residence time of the fluid inside the pellet and the efficiency of reaction is increased with a higher generation of product or it is decreased due to encapsulating effect of the reaction product into disconnected pores\(^\text{11}\). Therefore the clarification of fluid flow and heat transfer phenomena within the pores of the agglomerates offers opportunities to enhance the efficiency of the unit operation of various industrial processes based on self-reducing agglomerate\(^\text{10,11}\). In this context, the serial sectioning technique combined with chemical analysis and thermodynamic calculations are considered of low cost and useful tools to address detailed study on self-reducing agglomerates. The serial sectioning technique consists of three-dimensional numerical reconstruction of the microstructure from consecutive images obtained by optical microscope, for the superficial thin sections of the sample, removed by polishing\(^\text{5,12,13}\). This technique includes sequential steps as following: mechanical or chemical polishing of the sample surface; Vickers microhardness measurement; two-dimensional images acquisition in optical microscope; images alignment and three-dimensional reconstruction and visualization of the 2-D plans\(^\text{14}\). For each polishing step, the same amount of material is removed and others microhardness measurements are performed\(^\text{15}\). In this work the serial sectioning methodology was performed using the free software NIH ImageJ\(^\text{16}\) in order to visualize the inner structure of the pores present in the self-reducing pellets. The next step consists to measure the average tortuosity and the porosity values which plays important role on the mechanical strength of the agglomerates and heat and on mass transfer phenomena, which in turn, determines the efficiency of the processing technology based on the self reducing agglomerates\(^\text{10}\). When the self-reducing of the pellet begins, the reducing gases formed in the inner of the agglomerate, in contact with the reagent, make the chemical reactions according Equations 1, 2, 3 and 4, been the controller of the kinetic reduction process.

\[
\begin{align*}
3\text{Fe}_2\text{O}_3(\text{s}) + \text{CO}_2(\text{g}) &\leftrightarrow 2\text{Fe}_3\text{O}_4(\text{l,s}) + \text{CO}_2(\text{g}) \\
\text{C(\text{s})} + \text{CO}_2(\text{g}) &\rightarrow 2\text{CO}(\text{g}) \\
\text{Fe}_3\text{O}_4(\text{l,s}) + \text{CO}_2(\text{g}) &\leftrightarrow 3\text{FeO}(\text{l,s}) + \text{CO}_2(\text{g}) \\
\text{FeO}(\text{s}) + \text{CO}_2(\text{g}) &\leftrightarrow \text{Fe}(\text{s}) + \text{CO}_2(\text{g})
\end{align*}
\]

The overall mechanism is believed to be controlled by the rate of solution loss reaction (Equation 2)\(^\text{5}\). A rate equation largely used to model the kinetics of these processes is given by Equation 5. This kinetic reaction rate equation depends on the inner structure of the agglomerates such as contact areas, porosities and tortuosity, besides the gas diffusivities and activated sites for the reaction taking place\(^\text{17}\).

\[
R_{\text{CO}_2} = \left( \frac{\rho P_{\text{CO}_2}}{M_{\text{CO}_2}} \right) \left[ \frac{d_{\text{par}} \varphi}{A_{\text{par}} D_\text{bulk}^{\text{eff}} k_{\text{CO}_2}^{\text{bulk}}} + \frac{1}{\varphi_{\text{kinetic}} k_{\text{CO}_2}} \right]^{-1}
\]

In Equation 5, the parameter \(A\) is the superficial area of the particle that is function of the form factor and of the volumetric fraction as shown in Equation 6; and \(\eta_{\text{ave}}\), is a diffusion term that involves the tortuosity and porosity parameters, respectively, \(\tau\) and \(\xi\) according Equations 7, 8 and 9.

\[
A_{\text{par}} = 6(\varepsilon f / (d\varphi))
\]

\[
\eta_{\text{ave}} = \left( \frac{1}{\tanh((3\Theta) - 3\Theta)} \right)
\]

\[
\Theta = \frac{1}{6} (d\varphi_{\text{pch}}) \left( \frac{D_{\text{CO}_2}^{\text{eff}}}{v_{\text{CO}_2}} \right)
\]

\[
D_{\text{CO}_2}^{\text{eff}} = \frac{\tau_{\text{pch}}}{\tau_{\text{pch}}} (\frac{k_{\text{CO}_2}}{D_{\text{CO}_2}^{\text{bulk}}})
\]

In this study, the porosity obtained by the serial sectioning methodology was compared with traditional pycnometric technique. In addition, the agglomerate was characterized by chemical analysis using the Inductively Coupled Plasma Atomic Emission Spectroscopy (ICP), Scanning Electron Microscopy, Optical Microscopy and X-ray diffraction to recognize the present phases and additional discussions to compare the techniques. All the samples were prepared with pellet size compatible with the traditional pellets largely used in the reduction processes which would be easily applied for all proposed self-reducing processes.

2. Experimental Procedure

2.1. Chemical characterization

The chemical composition of the pellet was identified using the Inductively Coupled Plasma Atomic Emission Spectroscopy technique (ICPAES). In order to compare the results obtained, a chemical analysis was carried out by EDS (Energy Dispersive X-ray Detector) coupled in a Scanning Electron Microscope Zeiss EVO MA 10 for different regions of the sample. To recognize the phases presenting in the pellet made of electric arc furnace dust, the X-ray diffraction analysis was performed for the pulverized sample. From the software Match 1.10a the main phases of the sample were recognizing, using the database PDF (Powder Diffraction File). To confirm the composition homogeneity in whole volume of the pellet, the sample was cut into four equal parts and each part was pulverized and analyzed by X-ray diffraction. The X-ray diffractometer XRD-6000 Shimazu worked with a current intensity of 30 mA, 30kV voltage and the angle scan range was from 30° to 100° with a step of 0.02°. The composition of oxygen and others elements obtained by X-ray diffraction were calculated using mass balance from all phases detected. The Inductively Coupled Plasma Atomic Emission Spectroscopy (ICPAES) technique is considered more reliable, but, it was not possible to detect...
oxygen and carbon. Due the high amount of carbon found in X-ray and EDS not be reliable, the calculations for others elements detected were rearranged using calculation basis on 15% of carbon, a value commonly used in production of pellets with this kind of raw material.18

2.2. Three-dimensional microstructural characterization

The metallurgical preparation consisted to ground the sample in SiC paper (120-2500 grit) and after this, the polishing was performed with the dry sample (always that an image was acquired), using SiC paper with 4000 grit in a sanding machine PLOZED. It was not possible to perform the polishing with chemical suspension, because this could damage agglomerate structure of the sample and destroy the pores of the surface. In order to keep the same reference flame and return at the same position the microhardness was measured near the area to be analyzed. This procedure was repeated for each new polishing and image acquisition for 85 sequential images. Microhardness Vickers measurements were carried out in a Shmadzer model binocular microscope composed by lens of 0.65M 40x, and the penetration time of the indenter was fifteen seconds and the hardness required was 2.94N. The digital images were acquired with a 5X magnification in an optical microscope Leitz model Metallovert, using the software PixeLink Capture to acquire the images (dimensions 1280 × 1024 pixels). At each polishing step carried out, the spacing between two slices was computed from the reducing of diagonals of Vickers marks, observed in two sequential images. The thickness (Δh) of material removed at each polishing performed, is computed from the difference between mean values of Vickers diagonals before and after polishing (Dl − Dr) by Equation 10:

\[
\Delta h = \left( \frac{D_l - D_r}{2 \tan(\varphi/2)} \right)
\]

Where, \( \varphi \) is the angle between opposite faces in square-based pyramid with value of 136°. The three-dimensional visualization was processed using the freeware software NIH Image J, following the sequential steps: i) segmentation of images: this step is used to discern the pores, by pixels differentiation, of the other phases (pattern); ii) alignment of the 2-D planar sections using the available plugin “StackReg” developed by Philippe Thévenaz, from the Biomedical Imaging Group of the École Polytechnique Fédérale de Lausanne. The plug-in was based on four steps that were used in this work: translation, rigid body, rotation and affine. This alignment is necessary to ensure that the pore analyzed in the previous image is the same one which is visualized in the posterior image. The alignment process finishes only when the images are perfectly aligned to each other. Sometimes, the alignment was necessary to be performed manually using the plugin “Align Slice”20, and, iii) volume renderization using “Volume Viewer”21 plugin to allow a more “solid” visualization of the microstructure and after removing the pattern, it was possible to visualize the porous inner structure of the reconstructed volume. The amount of material removed after each polishing step was about 2.82 μm; according to the literature the amount of material removed is accepted until 3 μm, since above this value, some important information could be lost. Considering each image acquired with the same thickness than the material removed, and that, eighty five images were obtained, so, the computed volume for the sample is 1485 × 1188 × 241 μm³. The tortuosity parameter was calculated for 20 different pores. In addition, the pore fraction was calculated for 5 fields with the software NIH ImageJ using the plugin “Voxel Counter”22 and the result was compared with the porosity obtained by the pycnometric technique and analysis of 2D digital images by SEM.

In order to compare the methodologies additional pycnometry measurements were carried out. The pycnometric technique is a comparative method between the relative densities of the solid and liquid.23 This method is based on ASTM B 311-93 standard and it is valid for metallic materials containing less than 2% porosity.23 Due to the porosity of the self-reducing pellet to be high, in this study it was necessary to coat its surface by applying a thin polymeric film in order to avoid the pores filled by the liquid and so, to invalidate the true value for the measureable weight. According to the Equations 11 and 12, it is possible to estimate the results for density and porosity of the pellet, respectively.

\[
\rho_a = \rho_l \left( \frac{P_a}{P_a - P_l} \right)
\]

\[
\text{pore}(\%) = \left( \frac{\rho_l - \rho_a}{\rho_a} \right) \times 100
\]

Where, \( P_a \) is the weight of dry sample (kg.m/s²), \( P_l \) is the measurable weight of the sample (kg.m/s²), \( \rho_l \) is the liquid density (g/m³), \( \rho_a \) is the theoretical density (g/m³) and \( \rho_a \) is the apparent density (g/m³). The theoretical density can be determined by a theoretical calculation. In this work, the theoretical density was obtained using mixture rule based on the volume fraction of each phase identified by using the software Match. The density and pores fraction were calculated by 5 different pellets through the pycnometer. The liquid used was water and for each pellet, the measurements for measurable and true weights were accomplished five times. In addition, a 2D image analysis was carried out to compare the accuracy for the 3D method. This was carried out using the software NIH ImageJ, with the available resource Voxel Counter. After the metallographic preparation of the sample, all the digital images obtained by SEM were processed and analyzed using the freeware for image analysis NIH Image J. The images were treated and converted to bitmap code and selected by the brightness cut off to identify the pores. The pore fraction in 2-D was measured for five random areas.

3. Results and Discussions

3.1. Results of chemical characterization

In order to estimate the quality of the pellet produced with EAF dust, the elements presents in the pellet were identified and quantified by ICPAES analysis. To compare and ensure the accuracy of the results obtained by ICPAES
technique, EDS and X-ray analysis were carried out for different areas of the sample and the results using EDS method is presented in Figure 1, while x-ray spectra is shown in Figure 2. Both techniques showed a high value of carbon, and as it was explained (see section 2.1) the carbon fraction was estimated on 15% and the fraction of others elements were rearranged according to this new argument adopted. The EDS results are obtained using the averaged value of 5 random regions of the sample. In Table 1, according to the results of ICPAES, EDS and X-ray analysis, a high concentration of iron was identified as Fe$^{3+}$ or Fe$^{2+}$ ions and also like neutral element$^{23}$. In addition, some elements in low concentration as 6.30% calcium, 3.12%, manganese, and 3.06% magnesium, detected by ICPAES could be confused with the crystallographic structure of silicon found in high amount as SiO$_2$ phase by X-ray diffraction. The zinc concentration of 10.63% in EDS technique, was identified as ZnO and associated with iron as franklinite, according to the results obtained by X-ray diffraction. These results are in agreement with previous works reported at the literatures$^{24,25}$. The quantity of oxygen and carbon detected by EDS technique are not reliable since a few fields of the sample were analyzed however, mass balance of these elements were used to estimate the elements of inaccurate determination. The results obtained by X-ray diffraction show the major elements are carbon, oxygen (by mass balance), zinc and iron. As presented in Figures 1, 2, this study confirms the range of the main components presented in the self-reducing agglomerates of EAF dust obtained by previous works$^{6-8,24-27}$ besides the higher concentrations of zinc and iron based phases. Figure 2 shows the mean result for spectra obtained for each part of the pellet using the software Match v.1.0, and the composition of each phase calculated by the software. The X-ray diffraction results of the different regions of the sample were similar indicating the homogeneity of pellet composition. The blue spectrum is the experimental result and the color peaks are the spectra calculated based on available PDF database and its mass composition of the phases estimated by the software based on Rietveld method. The software also estimates the $R_p$ parameter that is a measure of confidence on quantifying the phases by the Rietveld method. Its value is considered in acceptable region between 15 and 30%$^{28}$. From the results of the composition of the phases, the calculation was performed on mass fraction presented in Figure 2 for each component through a simple mass balance. It is possible to observe the large amount of magnetite, franklinite, zincite, wustite and quartz. According to literatures$^{5-9,26,27}$, magnetite is commonly detected in high amount in electric arc furnace

![Figure 1. EDS spectra for random area of the self-agglomerate pellet of EAF dust. *Quantify of carbon and oxygen is not reliable.](image1)

![Figure 2. Average spectra for the 4 area of the self-reducing pellet produced with EAF dust. *Quantify of carbon (graphite) is not reliable.](image2)
dust. This characteristic of the EAF dust is confirmed in this study, as observed in Figures 1, 2. A summary of the average elemental analysis obtained by different techniques is presented in Table 1 for self-reducing pellets of EAF dust. In this kind of material, the carbon content is higher than the necessary amount for the reduction of the oxides in self-reducing pellets of EAF (usually until 15%) and contributes to dilute the large zinc amount present in the EAF dust and enable the production of agglomerates suitable to be used as raw materials for processes based on self-reducing technology.\(^{17,24-27}\)

### 3.2. Results for microstructural characterization

#### 3.2.1. Two-dimensional results

The analysis of the digital images obtained by SEM allowed to identify the pores present in the surface of the sample and thus, quantifying the pores fraction for posterior comparison with the 3D reconstruction technique. Figure 3 shows the images of a random area of the sample where the contrast of the pores in the images is poor and thus the pixel selection is performed to get better contrast using the software NIH ImageJ. After the segmentation step the resource termed analyze particles, available in the software, allowed the pore fractions calculations. The analysis of the different areas indicates that there was a large variation on the results of pores fraction in the different areas and this can be explained by the strong anisotropy of the material. It was observed that the distribution of pores in the pellet is higher in the central region of the agglomerates.\(^{3}\) The average porosity found for the different areas was 22.4%. In contrast, Wagner et al.\(^ {3} \) measured the pore fraction of pellets of iron ore shows slightly higher values. It is worth to mention that the morphology and spatial distribution of the pores in agglomerates play a relevant role in compressive strength and transport phenomena.\(^ {29,30} \) Therefore, a method able to furnish accurate results is of fundamental importance. This

| Elements   | ICPAES Composition (wt. (%)) | EDS Composition (wt. (%)) | X-Ray Composition (wt. (%)) |
|------------|-----------------------------|---------------------------|-----------------------------|
| Aluminium  | 0.24                        | 0.25                      | 0.00                        |
| Calcium    | 6.30                        | 3.12                      | 0.00                        |
| Cobalt     | 0.02                        | 0.00                      | 0.00                        |
| Chrome     | 0.33                        | 0.00                      | 0.00                        |
| Iron       | 53.64                       | 41.78                     | 34.82                       |
| Potassium  | 1.57                        | 0.48                      | 0.00                        |
| Magnesium  | 3.06                        | 1.41                      | 0.00                        |
| Manganese  | 3.12                        | 0.99                      | 0.00                        |
| Nickel     | 0.12                        | 0.00                      | 0.00                        |
| Silicon    | 1.98                        | 0.78                      | 10.75                       |
| Zinc       | 14.61                       | 10.63                     | 13.86                       |
| Carbon     | 15.00                       | 15.00                     | 15.00                       |
| Oxygen     | 0.00                        | 25.00                     | 25.56                       |
| Molibdenium| 0.00                        | 0.37                      | 0.00                        |
| Chlore     | 0.00                        | 0.19                      | 0.00                        |
| Total      | 100.00                      | 100.00                    | 100.00                      |

![Figure 3](image3.png)

**Figure 3.** Selection of the pixels for the porosity identification in a random area of 2D image.

![Figure 4](image4.png)

**Figure 4.** Sequence of 2D images treatment to carry out the 3D reconstruction of the porosities.
work aims to furnish comparative methodologies analysis to estimate inner parameter for modeling inner phenomena.

3.2.2. Three-dimensional results

In order to construct the 3D images, Figures 4 and 5 are used. Figure 4 shows the pixels selection while Figure 5 shows the sequence and alignment of 2D images for 3D reconstruction. At each polishing step, the material amount removed between two sequential images showed some negligible discrepancy in the values. This happens because the material is anisotropic and some parts are softer than others and careful hardness mark is important to assure same amount of removed material. Equation 10 allows the calculation the thickness of the material removed in each polishing cycle and the result found was 2.82 μm and assumed for all steps. To remove false pore caused by image noise which appeared in the pictures, it was applied some resources available in the software NIH ImageJ *(Filter plus and threshold)*. This step, called segmentation, consists of distinguishing the interesting phase of the matrix by pixels differentiation. Figure 4, from the left to right, the first picture is the original photo of a random area of the sample; the second photo is the original photo after the treatment of pixels using the resources *smooth* to round the bounds of the phases; *enhance contrast and filter plus* and *fast filter* to remove the noises of the images; *contrast and brightness* to distinguish and emphasize the contrast between matrix and pores; and in the third photo the image was changed for white (matrix) and black (pores), after to apply the *threshold* resource of the software *NIH ImageJ*. This treatment was done for the eighty and five 2-D images after to build a stack of images. Due to the rotation and displacement possibly generated during image acquisition, it was necessary to use the plug-in *StackReg*. When the pictures showed a high distortion compared to other it was necessary to perform the alignment manually using the *Align Slice* plug-in. Figure 5 presents an example of the alignment for six sequential images (from (a) until (f)) of a random field. In Figure 5 it is easily noted a dark region formed by pores closer to each other caused by microhardness mark that must be removed from the image analysis in order to avoid over prediction of the pore volume fractions. In addition, it is possible to note that some pores disappear away from the second to third image, and new pores arise in the fourth until sixth image. After the alignment of the images the fields were reconstructed using the resource *Volume Viewer*. Much of the presents pores visualized in Figure 6 were classified like micropores and it was not possible to

![Figure 5. Alignment of the sequence from (a) until (f) of 2D sectioning for the 3D reconstruction of the self-reducing pellet.](image)

![Figure 6. Results of the 3D reconstruction of the pores in a portion of the self-reducing pellet.](image)
visualize the connectivity after the polishing. Figures 6a-c, show the reconstructed images of some fields without the matrix. Some pores showed an irregular structure, but the majority of the pores show a nearly cylindrical geometry. The micropores are nearly spherical shapes. The majority of isolated micropores arise inside some grains and phases. This usually happens during the agglomeration process, when the growth of the grains causes the detachment between the pores and the spherical geometry of these pores that remaining in the grain\textsuperscript{20,26}. Figure 7 is an example of 3D image reconstruction field with details to visualize the morphology of the pores present. Isolated pores and several distorted shapes were observed.

3.3. Predictions of tortuosity and porosity

The tortuosity and porosity parameters in the pellets are extremely important for the kinetics of reduction reactions. Higher tortuosity values is expected to increase the turbulence and the residence time of the gas (CO) inside of pellet\textsuperscript{29}. The tortuosity is defined mathematically by Equation 13.

\[
\tau = \left( \frac{L_v}{L_l} \right)^2
\]  

(13)

In this study the tortuosity parameter was calculated for 20 pores reconstructed in the sectioning area of the sample. Figure 8 allows to observe how this calculation was carried out using the \textit{ImageJ} software and the legend attached to the figure displays the values of the real ($L_v$) and linear ($L_l$) lengths in μm and the respective tortuosities ($\tau$) for the 20 different pores reconstructed. The average value for tortuosity was 0.82. It is a value close to 1 that suggests a nearly cylindrical geometry shape of the large pores. In general, the pores show real path length longer than the straight ones in the direction of fluid flow. Excluding the 7\textsuperscript{th} pore measured, the tortuosity value found was about 0.80. The pores 2, 3, 9 and 12 had the larger tortuosity, 1.00, indicating cylindrical structure. The porosity was calculated by plug-in \textit{Voxel Counter}. This step was performed by the ratio between the voxels of the pores present in the analyzed field and voxels of the entire field. The porosity was calculated for 5 different fields and the superior pores distribution was 48.81\%, and the inferior, 32.54\%. This difference is explained due the material heterogeneity and the porosity increases from the surface to the center of the pellet\textsuperscript{3}. In addition, the porosity was calculated by the pycnometric technique, which is assumed as the best method for volumetric bulk measurements, for 5 different pellets. With Equations 2 and 3, the apparent density and pore fractions were estimated assuming that the theoretical density ($\rho_t$), the weight of dry pellet ($P$), apparent weight ($P_a$), apparent porosity ($\rho_a$) and fraction of pores for 5 different self-reducing pellets of EAF were calculated. The porosity range was from 39\% to 43\% showing that the 3D sectioning technique is compatible with pycnometry. Figure 9 shows...
the comparative results obtained for the three techniques of porosity measurements considered in this work. The two-dimensional analysis of the images obtained in SEM shows the more discrepant value. Probably, it occurs due to 3D features of pores in the anisotropic materials. As it was observed, the three-dimensional reconstruction and pycnometry showed similar values due to their natural 3D features. Additionally, the analysis by two-dimensional images using SEM requires a large number of pictures of all surface of pellet and usually offers lower accuracy. Thus, this study recommended the 3D sectioning reconstruction as an accurate and feasible technique to detailed characterization of self-reducing pellets for pore measurement, inner structure and measurements of inner parameters of the pore such as tortuosity, connectivity and volume fractions of micropores and clusteres.

4. Conclusions

In this work the chemical and microstructural characterization of self-reducing pellets made of EAF dust was carried out. The ICPAES (Inductively Coupled Plasma Atomic Emission Spectroscopy) technique was considered suitable to recognize and to quantify the constituents present in the self-reducing pellet. The X-ray diffraction allowed to identify and to quantify the main phases of the pellet of EAF with typical values: magnetite (11.6%), wustite (14.05%), zincite (11.10%), quartz (19.93%), iron (5.50%) and carbon (15.00% estimated), commonly found in electric arc furnace dust [16,22]; and using mass balance it was possible to estimate the amount of oxygen present in oxides detected (25.56%). A large amount of quartz was detected by X-ray diffraction and perhaps, the silicon could have been confused by others elements in lower concentration with similar crystallographic structure. The three techniques allowed detecting a high amount of iron and zinc, commonly found in EAF dust. The three-dimensional reconstruction by serial sectioning technique was compared with those found by two-dimensional image analysis obtained in SEM and pycnometry technique. The average of results for porosity obtained for three-dimensional, pycnometry and two-dimensional techniques were, respectively, 41.61%, 41.53% and 23.41%. The 3-D method presented more accurate results, as expected, since the porosity is a parameter of typical three-dimensional feature especially for anisotropic materials. In addition, the 3-D technique has the advantage of visualizing the pore morphology and interconnectivity among them, an important reduction kinetic parameter of the self-reducing pellets. By observing the final reconstruction images it was found that the pores are close to cylindrical shape with low interconnectivity. In this work, the average value of tortuosity obtained for a typical self-reducing pellet of EAF dust was 0.84 that proposes a morphological structure close to cylindrical. In addition, the pycnometry allowed calculating the density of the pellet, 2.3342 g/cm³.

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Appendix

Nomenclature

| Symbol | Description |
|--------|-------------|
| Fe₂O₃ | magnetita |
| FeO   | wustita   |
| Fe₂O₃ | hematita |
| Fe    | iron       |
| ZnFe₂O₄ | zinc ferrite |
| ZnSiO₃ | zinc silicate |
| ZnO   | znicite |
| Fe⁺³ | ions iron III |
| Fe⁺² | ions iron II |
| τ     | pore tortuosity |
| Lv    | average length crossed by the fluid through the pores. |
| L     | Linear length in the direction of fluid flow. |
| D1    | mean diagonal obtained from images before and after polishing. |
| D2    | mean diagonal obtained from images before and after polishing. |
| φ     | angle formed between diagonal and height of microimpression geometry. |
| ρ₀    | apparent density of the pellet. |
| ρₜ    | theoretical density of the pellet. |
| pl    | fluid density (1 g/cm³). |
| P     | Weight of dried pellet. |
| Pₛ    | Apparent weight of the sample. |
| Rp    | confidence factor to quantify the phases by Rietveld method. |