Characterization of iron oxide waste scales obtained by rolling mill steel industry

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Abstract
Every year, the steelmaking industry produces millions of tons of slags resulting in pollution to the environment. Among the waste, secondary metals and scales rich in iron oxides are also thrown away. There is a need to treat the steel waste in a reasonably way to protect the environment and proposing new cheap technologies for producing advanced materials. In this study we report the morphological and structural characterization of waste scales generated during roll milling steel process at JSC “Arcelor Mittal Temirtau”. The raw slag and annealed at 1000 °C were measured by X-ray diffraction (XRD), scanning electron microscopy adapted with energy dispersive X-ray (SEM-EDX), magnetometry and Mössbauer Spectroscopy (MS). Fe and O were detected by EDX as main chemical elements and Si, S, Ca, Mg, C and Al as minimal elemental composition. XDR for the raw sample revealed α-Fe2O3 (hematite) and Fe3O4 (magnetite) as principal and secondary phase, respectively; whereas monophasic α-Fe2O3 is detected for the scales annealed at 1000 °C. Magnetometry measurements show the Verwey transition for the raw sample and the Morin transition for the annealed at 1000 °C; those are fingerprints for the presence of magnetite and hematite, respectively. MS measurements for the raw sample consist of 6 small peaks of absorption and a broad two-lines absorption peak in the central part. The doublets are associated to the hyperfine parameters belonging to wustite. Magnetite is related to the hyperfine parameters for two sextets in octahedral Fe2.5+ and tetrahedral Fe3+ sites and a small sextet that resembles the Mössbauer parameters of α-Fe2O3. Only a well crystallized and weakly ferromagnetic sextet confirm the presence of α-Fe2O3 phase for the sample annealed at 1000 °C due to thermal oxidation.

Keywords Iron oxide · Steel slag · Steel scales · Rolling mill steel · Mössbauer spectroscopy
1 Introduction

During the industrial production of iron steel, tons of iron-rich scales are produced as waste materials [1]. Since the waste contains large amounts of iron oxides, heavy metals and other different contaminants, it affects the environment when disposed in landfills because it releases toxic substances in soil and groundwater [2]. Thus, steel wastes create pollution to the environment. If these resources are reasonably treated, then not only the environment would be protected and disposal costs reduced, but also proposing new cheap technology for producing advanced materials [3]. Some attempts in re-using waste materials from steel industry includes iron oxide pigments, new metal pieces, manufacturing equipment, chemical tankers, automotive and airplane components [4, 5].

For the recycling, the steel waste should be previously treated in different ways, such as grinding, pressing, bathing, vitrifying, annealing, and other related procedures under various conditions of pressure, atmosphere and temperature [1, 6, 7]. For example, annealing promotes the oxidation of iron such as hematite (α-Fe₂O₃) and magnetite (Fe₃O₄) and then the iron components can be separated magnetically. In this way, it is very important to know the composition and characteristics of the waste materials before attempting to recycle them [1] [8–11]. Carlini et al. have observed that by regulating the number of iron ions in the body and limiting the crystallization process, it is possible to obtain materials with conductivity less than 1000 Ω [3]. Also, it has been reported that the formation of nano- or micro-scale crystals gives iron oxide certain outstanding properties, such as electrical conductivity and ferromagnetism [6]. Moreover, the characteristics of pigments obtained from recycled iron oxides are greatly affected by the grinding, milling and heat treatment processes and also by hematite content [2]. In addition, oxide scales obtained during rolling have different magnetic properties after magnetic separation [6].

Therefore, for a correct recycling and application of industrial steel waste, it is necessary to have good information on its composition, characteristics and properties. In this work, we report the characterization of iron oxide waste scales obtained by rolling mill steel industry. The sample was obtained during the mechanical cleaning of the surface of hot-rolled steel strip in the production process. This is a promising prime material since its price is much cheaper than hematite powder obtained during the pyrolysis of spent pickling solutions.

2 Experimental

The scales formed on the surface of steel sheets during the hot rolling process (T ~ 700-1200 °C) of steelmaking were collected [12]. It consists of mill scales of wustite (FeO), magnetite (Fe₃O₄) and hematite (α-Fe₂O₃). The collected sample was also annealed at 1000 °C in air. The morphology of the samples was inspected through a scanning electron microscope (SEM) Philips XL-30 equipped with an EDX X-MAX (80 mm²) Oxford Instruments Inc. and acceleration voltages of 20-30 kV. The X-ray diffraction (XRD) were taken in a Rigaku diffractometer with Cu-Kα radiation λ = 1.5409 Å with 20° < 2θ < 90°, 0.01 step and scanning speed of 0.01°/s. The current was 200 mA. Mössbauer spectroscopy (MS) measurements at room temperature were carried out using a Mössbauer spectrometer in transmission geometry. A ⁵⁷Co source in Rh matrix with 50 mCi of nominal activity moving at constant acceleration. For the calibration measurement, a thin foil α-Fe standard
sample was used. The fitting process was achieved applying least square theory with Lorentzian profiles in the WinNormos software. The magnetic measurements were performed using a DC-MPMS-SQUID magnetometer (Quantum Design Inc.) in zero-field cooling (ZFC) and field cooling (FC) modes. The magnetization response as a function of the temperature was collected from 5 to 400 K and applying external magnetic fields between 100 Oe to 10 kOe. The magnetization responses as a function of an external applied magnetic field were also measured in the range ±10 kOe and at different temperatures between 5 to 400 K.

3 Results and discussion

Figure 1 displays the SEM micrographs and the EDX spectra of the raw scales and after annealing at 1000 °C. The SEM micrographs of the raw sample (see Fig. 1(a)) reveal that the morphology of the scales has lamellar shapes of different sizes, ranging from millimeters to several centimeters. They are flat and spread without sticking together. Figure 1b) is a zoomed view which reveals that the scales are composed of grains with planar fractured faces and stuck together. The faces of the grains are smooth suggesting good crystallinity although the grain boundary is not appreciable. Figures 1 (c and d) show the micrograph of the samples after annealing at 1000 °C. The high temperature provokes that the sample becomes a lump. The scale morphology is lost and the grains tend to stick together; the faceted grains decrease their size suggesting melting in the grain interfaces. In Fig. 1e) the EDX mapping spectra for the raw sample shows that the main elemental components are Fe and O, suggesting the presence of iron oxides as main mineralogical phases. Note that some spots of Ca are spread over the samples. Table 1 lists the quantification of the main elements obtained by EDX. Ca may proceed for the non-reacted calcium carbonate added during the steelmaking process [13]. In addition, other minor elements, such as Si and Al are found and might belong to silicates and aluminosilicates coming from soil dust present in the collection zone. Figure 1f) shows the EDX mapping of the scales after annealing at 1000 °C. Similarly, to the previous case, Fe and O are the main compositional elements. Note that in addition to the Si, Al and Ca assumed from silicates from soil dust, other minor components, such as Mg, Cr and C appear. Carbon could come from contaminants from the furnace processing, Mg might come from remaining unreacted compounds during processing the iron steel, while S was added during steelmaking to improve machinability [14].

Figure 2 shows the X-ray diffraction for the collected scales and for the samples annealed at 1000 °C. The main phase for the raw sample is α-Fe₂O₃ (hematite). Some tiny peaks reflections are also found between 25° and 33°, indicated in bold filled squares, which could be related to some silicates compounds according to the EDX results above. The diffraction pattern of the sample obtained after annealing at 1000 °C shows only hematite phase. Note that some peaks become sharper, indicating that the crystallization of the hematite has improved. Miller indexes, angles, full width at the half maximum (FWHM), and crystallite sizes obtained from the XRD are listed in Table 2. FWHM tends to decrease its values meanwhile the grain size increases due to thermal heating. This indicates that hematite crystals grow by thermal induction.

Note that at higher angles than 75°, the peaks belonging to the (220), (036), (128), (0 0 12), and (226) reflections appear (enclosed by the dash squares). This also confirms the improvement in crystallization of the hematite phase due to the oxidation of other possible
Fig. 1 SEM micrographs and EDX analysis of iron oxide waste scale obtained by rolling mill steel industry and after annealed at 1000 °C. (a) and (b) micrographs of the collected scale samples and (c), (d) micrographs for the annealed sample at 1000 °C. (e) and (f) EDX for raw sample and after annealing at 1000 °C.

Table 1 Elemental quantification in percentage weight (wt%) obtained by EDX analysis for iron oxide waste scales obtained by rolling mill steel industry and after annealing at 1000 °C.

| Sample   | O      | Fe    | Si    | C    | S     | Ca    | Al    | Cr    | Mg   |
|----------|--------|-------|-------|------|-------|-------|-------|-------|------|
| Raw      | 24.26  | 72.9  | 0.91  | 0.59 | 0.71  | 0.62  | 0.12  |       |      |
| 1000 °C  | 22.83  | 65.89 | 1.01  | 3.39 | 0.17  | 0.3   | 1.99 ± 0.13 | 1.83 ± 0.02 |
minor iron oxides components. In this case, the presence of those peaks indicate the complete oxidation of magnetite [15], as detected by the other techniques below.

The temperature dependence of the magnetization (M(T)) curves for the raw scales obtained during rolling mill process of steel industry is given in Fig. 3 (a). The M(T) curves show decrease of irreversibility in ZFC and FC as the applied external field is increased between 100 Oe to 5 kOe. Increasing the applied external field provokes the alignment of more magnetic domains and resulting in the gaining of magnetization. Moreover, at each external applied magnetic field, the M(T) responses show the Verwey transition at T ~ 124 K which is the usual fingerprint to identify magnetite [16]. Note that the XRD analysis above detects also hematite for this sample. However, since the magnetization values for hematite are very low, it could be screened by that for magnetite and others magnetic phases.

The M(T) responses for the scales annealed at 1000 °C are given in Fig. 3 (b). They show low magnetization values compared to the raw sample (see Fig. 3 (a)). The Verwey transition is not detected anymore and instead, the Morin transition remarkably appears. The Morin transition is the fingerprint signal for presence of hematite [17]. It is not screened at higher external fields meaning that any other ferromagnetic or anti-ferromagnetic domains are not present. Nevertheless, there is a paramagnetic signal, which becomes noticeable when increasing the external magnetic field. These results confirm that magnetite oxidizes by thermal induction after annealing at 1000 °C into hematite [15].

Figure 3 (c) shows the applied magnetic field dependence of the magnetization M(H) for the raw sample. It reveals hysteresis, typical of soft ferromagnetic material. The inset in Fig. 3 (c) displays the remanence magnetization values that decays with increasing temperature. However, the variation of remanence values does not show a clear tendency between 150 and 275 K. Similar behaviour is observed in coercivity values (see inset in Fig. 3 (c) for raw samples). The decreases in remanence and coercivity values is due to the competition between thermal and magnetic energy in magnetic domains. That is, at low temperatures, thermal energy decreases, making spins stand fixed (high coercivity) resulting in more difficulty to reorient them. However, the variation in both remanence and coercivity in the temperature range between 150 and 175 K does not follow a defined trend. This indicates an incoherent spin rotation mode which must be caused by competition between
the magnetic moments of the different components of the sample [18]. This effect does not happen in scales obtained by rolling mill annealed at 1000 °C because the major component is hematite, as it is discussed next.

Figure 3 (d) shows the applied field dependence of the magnetization (M(H)) loops for the annealed sample. They reveal hysteresis due to the presence of hematite coexisting with paramagnetic domains. Note that the hysteresis loops near the Morin temperature at T ~ 275 K show deformed hysteresis. That is, near zero applied field, the hysteresis curves are less wide than with increasing external magnetic field (see bottom inset plots). This is due to the presence of weak antiferromagnetic and ferromagnetic domains at this temperature [18]. Note also that, according to the top inset plots, the remanence and coercivity values increase slightly above 250 K, this is due to the transition of hematite from antiferromagnetic to weak ferromagnetic.

The RT Mössbauer spectra of the raw scales obtained by rolling mill steel industry and the annealed sample at 1000 °C are shown in Fig. 4. The spectra were fitted using

| Sample   | (hkl)  | 2θ (°) | FWHM (°) | Size (nm) |
|----------|--------|--------|----------|-----------|
| Raw      | (012)  | 24.11  | 0.28     | 29        |
|          | (104)  | 33.11  | 0.18     | 46        |
|          | (110)  | 35.55  | 0.47     | 18        |
|          | (006)  | 39.24  | 0.21     | 40        |
|          | (113)  | 40.8   | 0.17     | 49        |
|          | (024)  | 49.42  | 0.27     | 32        |
|          | (116)  | 53.97  | 0.26     | 40        |
|          | (018)  | 57.51  | 0.27     | 33        |
|          | (214)  | 62.45  | 0.34     | 27        |
|          | (300)  | 64     | 0.36     | 26        |
|          | (1 0 10)| 71.87  | 0.4      | 24        |
|          | (0 2 10)| 82.83  | 0.39     | 27        |
| 1000 °C  | (012)  | 24.12  | 0.15     | 54        |
|          | (104)  | 33.14  | 0.15     | 55        |
|          | (110)  | 35.59  | 0.14     | 59        |
|          | (006)  | 39.27  | 0.17     | 49        |
|          | (113)  | 40.85  | 0.19     | 44        |
|          | (024)  | 49.46  | 0.22     | 39        |
|          | (116)  | 54.11  | 0.22     | 40        |
|          | (018)  | 57.62  | 0.27     | 33        |
|          | (214)  | 62.44  | 0.26     | 34        |
|          | (300)  | 64.01  | 0.28     | 33        |
|          | (208)  | 69.59  | 0.12     | 80        |
|          | (1 0 10)| 72.01  | 0.33     | 29        |
|          | (220)  | 75.43  | 0.16     | 62        |
|          | (036)  | 77.81  | 0.29     | 35        |
|          | (128)  | 80.72  | 0.38     | 27        |
|          | (0 2 10)| 82.96  | 0.2      | 52        |
|          | (0 0 12)| 84.91  | 0.21     | 51        |
|          | (226)  | 88.71  | 0.43     | 26        |
Lorentzian profiles and the resulting hyperfine parameters are listed in Table 3. The Mössbauer spectrum for the raw milled scales, given in the top figure, presents six small absorption peaks and a broad one in the central part, these spectra were adjusted using three sextets and two doublets sites. The two doublets correspond to one paramagnetic dominant fraction with Fe$^{3+}$ site and relative absorption area (RAA) of 45.9%, and another paramagnetic fraction with Fe$^{2+}$ site and relative absorption 18.4%. The MS suggest that these two doublets correspond to the presence of wustite with hyperfine parameters $\delta_{\text{Fe}} = 0.86$ mm/s, $\Delta E_Q = 0.71$ mm/s and $\delta_{\text{Fe}} = 1.13$ mm/s, $\Delta E_Q = 0.63$ mm/s [19] which cannot be detected by XRD probably by its low quantity. Note that the quantity of Si is about 1% so it is no possible to associate it to any iron-silicate minerals. The particles sizes in Fig. 1(a) and (b) are not in the nanometer order and the M(H) curves show hysteresis loops in Fig. 3 meaning that the doublets do not correspond to superparamagnetic domains [20]. The two sextets are related to the octahedral Fe$^{2.5+}$ and tetrahedral Fe$^{3+}$ sites with hyperfine parameters $\delta_{\text{Fe}} = 0.86$ mm/s, $2\varepsilon = 0.04$, $B_{\text{hf}} = 45.9$ T and 21.6% RAA, and $\delta_{\text{Fe}} = 0.27$ mm/s, $B_{\text{hf}} = 45.9$ T and (RAA) of 12.6%, respectively. Since the magnetic measurements above detected the Verwey transition (see Fig.3 (a)), these hyperfine parameters correspond to magnetite [16, 21], which is in a medium absorption fraction content. Note also that the rate $\text{RAA(Fe}^{2.5+})/\text{RAA(Fe}^{3+})$ results in 1.7, indicating that magnetite is not in a stoichiometric ratio that means that it is partially substituted or oxidized, probably due to environmental interaction. The tiny sextet

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**Fig. 3** Magnetic responses of iron oxide waste scales obtained by rolling mill steel industry (a) M(T), (c) M(H) for the raw sample, (c) M(T) and (d) M(H) for the sample annealed at 1000 °C
with hyperfine parameters of $\delta_{Fe} = 0.32$ mm/s, $2\varepsilon = -0.23$, $B_{hf} = 51.7$ T and (RAA) of 1.5% is related to hematite [22] as detected by the XRD above. However, in contrast to the XRD analysis above which detects hematite as major mineral component and similarly to the magnetometry analysis above, the tiny sextet in the MS corresponding to hematite must be screened by the magnetite sub-spectra.

Regarding the scales annealed at 1000 °C, in agreement to the analysis by the other techniques above, which reveal the presence of monophasic hematite; the resultant MS spectrum consists of one sextet with hyperfine parameters $\delta_{Fe} = 0.37$ mm/s, $2\varepsilon = -0.21$, $B_{hf} = 51.8$ T and (RAA) of 100% which correspond to hematite. This confirms the thermal oxidation from wustite to magnetite and to hematite improving crystallization [15].
4 Conclusions

Iron oxide waste scales obtained by rolling mill steel industry was annealed at 1000 °C and characterized by different techniques. Fe and O are the major elemental composition and Si, S, Mg, Cr and Ca are the minimal components for the raw and annealed at 1000 °C samples indicating they consist mainly in iron oxides. α-Fe₂O₃ and Fe₃O₄ were the iron oxides detected for the raw sample scales, whereas α-Fe₂O₃ was the only iron oxide identified for the annealed sample at 1000 °C. Verwey transition, associated to magnetite content in the raw sample was registered in the M(T) curves under 100 Oe, 500 Oe, 1 kOe and 5 kOe external fields with high values of magnetization that screen the magnetic signal from hematite. The M(H) loops present hysteresis due to ferromagnetic domains. For the sample annealed at 1000 °C only hematite was registered by its Morin transition in the magnetic analysis. However, the hyperfine parameters confirm the presence of wustite and magnetite in non-stoequimetric state, and also hematite. Additionally, the MS reveals the formation of hematite for the annealed sample at 1000 °C due to oxidation of magnetite.

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