The effect of microstructure on the leaching behaviour of electric arc furnace (EAF) carbon steel slag

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A B S T R A C T

Electric arc furnace (EAF) slag could be exploited in several fields of application, such as land filling, road constructions and concrete production, since their physical properties are similar, or even better, than natural materials like gravel. Environmental concerns related to pollutants leaching (Ba, V, Cr, chlorides, fluorides, cyanides, etc.), are the primary hindrance limiting the effective reuse of such material. Thus, chemical and structural stability are fundamental requirements to be fulfilled, especially when the slag comes in contact with water. The slag microstructure has a non-negligible influence on metals concentration in leachate, even if the phases that react with water have not yet been clearly identified. In this study, different classes of carbon steel EAF slag were investigated in order to correlate their leaching behaviour with microstructural and crystallographic features. Fine particles were chosen to enhance the dissolution rate of the phases involved in the leaching process and to quantify the effects of the fine fraction on leachate concentration with respect to the bulk material. Qualitative tests at different liquid-to-solid (L/S) ratios were also performed to detect dissolution of the phases, which would otherwise be non-appreciable in standard conditions. The analyses also allowed for the identification of the phases responsible for toxic metal leaching (Ba, Cr, V), as well as those that were not involved in dissolution phenomena, thereby consolidating the results proposed elsewhere on slag stabilization.

Keywords:
Electric arc furnace (EAF) slag
Leaching behaviour
Microstructure
Fine fraction
Larnite
Gehlenite

1. Introduction

The growing interest in the use of steel slag as an alternative to traditional stone materials has prompted researchers to study in detail the chemical, physical and mechanical properties of these industrial by-products. Environmental concerns related to toxic metal leaching, are the primary hindrance limiting the effective reuse of such material, especially for slag from the electric steelmaking route. For this reason, several studies (Lind et al., 2001; Chaurand et al., 2007; van Zomeren et al., 2011; Engelsen et al., 2010, 2012) have investigated the leaching behaviour of slag aggregates, highlighting that such aggregates could potentially release dangerous chemical species (Ba, V, Cr, chlorides, fluorides, cyanides, etc.). Recently, strong correlations between leaching behaviour and the chemical composition of different classes of EAF slag have been highlighted and important considerations relating to the improvement of pollutant retaining substances were stated, concerning the standard test required by the EN 12457-2 regulation (Mombelli et al., 2014, 2016). Chemical composition plays a fundamental role and provides important indications on the leaching behaviour of a slag, since it determines, with cooling way, the slag microstructure (Tossavainen et al., 2007; Albertsson et al., 2013, 2014). Thus, microstructure has a non-negligible influence on metals...
concentration in leachate. As a consequence, one of the most important issues to solve is the identification of the mineralogical phases responsible for the release of toxic metals. In fact, not all the slag particle surface reacts with water, but only specific phases are involved in hydration and dissolution processes (Engström et al., 2013; Mombelli et al., 2016; Strandkvist et al., 2015). Several studies (Barella et al., 2012; Mombelli et al., 2012; Gellé et al., 2010; De Windt et al., 2011; Faellman, 2000; Engström et al., 2013; Strandkvist et al., 2015) indicate that calcium-rich crystalline phases (in particular larnite, calcium aluminates and calcium ferrite) are the most probable structural constituents involved in releasing phenomena, but which phases exactly react with water is not clearly identified. Moreover, the conditions under which the leachates are produced vary in relation to different physical parameters like granulometry, temperature or flow conditions, influencing the resulting concentrations of metals in the leachates (Faellman and Hartlén, 1996). Even if the different standard tests are carried out on bulk samples, a certain amount of fine particles is admitted, and these fine fractions negatively influence the test results, i.e. enhancing the leaching (Mazutani et al., 2006; Vlčková et al., 2011; Moser and Römcke, 2009). In fact, fine particles can lead to a variation of the pH of the eluate and consequently to a different leachability of pH sensitive constituents. For this reason, the European Standards usually specify that the material should not be finely ground and plan to limit particle size reduction in order to maintain the physical state of the waste. However, no indication about the minimum admitted size and amount of fine particles is indicated in the standards. In addition, since fine particles are usually employed as filler material in cementitious components, the leaching behaviour of such manufactures could be significantly changed by the filler weight fraction. Nevertheless, since the interaction between slag and water was experimentally demonstrated to be only cortical, interesting only the particle surface, the analysis of fine particles could allow to better identify the hydration and dissolution mechanisms that rule the pollutants leaching (Mombelli et al., 2014). Effectively, the increase in the surface-to-volume (S/V) ratio of fine particles, coupled with a high liquid-to-solid (L/S) ratio, favours the identification of dissolved phases. This is because it maintains the pH far from basic values, thus working in conditions a long way from the saturation limit, enhancing the dissolution rate as reported by Nicoleau et al. (2013).

In this study, different classes of carbon steel EAF slag powder originating from different steel productions (reinforced bar steel, high alloyed steel and quality steel) were investigated, in order to correlate their leaching behaviour with microstructural and crystallographic features. Fine particles were chosen to enhance the dissolution rate of the phases involved in the leaching process and to quantify the effects of the fine fraction on leachate concentration with respect to the bulk material. Moreover, qualitative tests at different L/S ratios were also performed to detect the dissolution of the phases, otherwise non-appreciable in standard conditions. The main goal of the present work is to identify which crystalline phases are responsible for the toxic metal leaching (Ba, Cr, V). The analyses have also allowed for the identification of the phases not involved in dissolution phenomena, thereby consolidating the results proposed elsewhere on slag stabilization by Mombelli et al. (2014; 2016). The results discussed in the present paper show quantitatively the relationship between leaching and the fraction of some mineral phases considered as key factors controlling the leaching behaviour of EAF slag.

2. Experimental procedure

2.1. Material

EAF slag were provided by different Italian and European electric steelworks and associated to different steel productions: reinforcing bar steel (group A, 7 samples), high-alloyed steel (group B, 4 samples) and quality steels (group C, 4 samples).

The investigated samples represent a selected subset from a larger sampling, previously analyzed by the same authors (Mombelli et al., 2016).

Fig. 1 – Particle size distribution of investigated slag powder.

The slag chemical composition was provided by ED-XRF analysis by means of Ametec Spectro Xepos spectrometer in He atmosphere on 5 g of powdered slag (Table 1).

The samples, in the form of powder obtained by ring mill grinding (particle size distribution in Fig. 1), were morphological and microstructural characterized by means of XRD and SEM analysis, and leaching tests were carried out in deionized water at pH 7, varying the liquid-to-solid ratio. To determine which phases were responsible for metals release, leaching tests on the polished section were also performed. Leaching tests results were correlated to chemical composition and crystalline phase fraction.

2.2. Crystallographic and microstructural characterizations

X-ray diffraction (XRD) data was collected using a Bruker D8 Advance diffractometer in a 0-θ configuration employing the Cu Kα radiation (λ = 1.54 Å) with a fixed divergence slit size 0.5° and a rotating sample stage. The samples were scanned between 10° and 80° (step size of 0.007°) with the Vantec detector. The qualitative analysis was performed with EVA software whereas a semi-quantitative analysis was performed by Crystal Match! Software, exploiting the Reference Intensity Ratio method (RIR-method) (de Wolff and Visser, 1988), calibrating the results on the base of a experimentally determined larnite fraction. Larnite fraction was determined through a selective dissolution in methanol-salicylic acid solution (Klemm and Skalny, 1977). 2.5 g of slag powder was mixed in 150 ml of methanol acidified with 10 g of salicylic acid. The solution was stirred for 2 h at 100 rpm and decanted for 15 min before filtering in a 0.45 μm vacuum system. The residual slag powder was dried at 105 °C for 1 h and weighed again (balance accuracy: 0.0001 g). The weight difference is a good estimation of larnite fraction.

Morphological and microstructural characterization was performed by Zeiss EVO50 Scanning Electron Microscopy (SEM) equipped with an Oxford Inca EDS probe. Slags were moulded in araldite-based resin, ground and polished.

2.3. Leaching test

Slag powder leaching behaviour was investigated by varying the L/S ratio (10, 100, 1000 l/kg). Weight loss, water pH and conductivity were measured after each test and the dried powders were investigated by XRD. The test consisted in immersing the slag powder in deionized water for 1 day,
keep shacked the PTFE bottle as recommended by EN 12457-2 standard. After 24 h, slags were removed and the water solution was filtered by 0.45 μm vacuum system and analyzed with a spectroscopic technique, i.e. ICP-OES (Induced Coupled Plasma-Optical Emission Spectroscopy, detection limit 50 g/l). The analysis focused on Al, Ba, Ca, Cr, Mg and V. In addition, the polished section of massive grains were leached in the same conditions as before and followed by a SEM investigation in order to detect which phases were dissolved or resist at a very high L/S ratio.

3. Results and discussion

3.1. Crystallographic and microstructural characterizations

In Table 2 the crystalline phases fraction is reported. The larnite percentage obtained by selective dissolution is also indicated. The XRD semi-quantitative analysis can be considered reliable since the difference between measured and calculated larnite fraction is negligible and at most less than 5%. Only sample #14 shows a large difference between calculated and measured larnite fraction. This could be due to the high sample heterogeneity.

In nearly every case the crystallographic analysis highlighted the occurrence of four phases¹ (Tossavainen et al., 2007; Shen et al., 2004; Shi, 2004; Luxan et al., 2000; Tsakiridis et al., 2008; Juckes, 2003; Reddy et al., 2006; Nicolae et al., 2007; Qian et al., 2002a,b; Manso et al., 2005; Geiseler, 1996; Wachsmuth et al., 1981): larnite, wustite, (Mg,Al,Fe,Cr) spinels, and gehlenite. Other phases, like brownmillerite, kirschsteinite or merwinite, were detected in specific samples, i.e.

3.2. Leaching test at 10 l/kg

First of all, to highlight the important effect of fine particles on leaching test, the comparison between leaching results in standard conditions (24 h mixing, L/S = 10 l/kg, deionized water pH 7) of powder and bulk samples was carried out. The bulk behaviour was reported by Mombelli et al. (2016).

The comparison highlighted the significant effect of particle size on dangerous metal concentration in the leachate samples #2, 8, 12 and 14. Only sample #8 was characterized by Ca-chromite, that appears as elongated lamellae (Fig. 2) inducing an intense diffraction peak at 2θ: 16° (Cabrera-Real et al., 2012; Albertsson et al., 2014).

By SEM-EDS analysis, the mapping of the toxic element (Ba, V, Cr) and their content in the different crystalline phases was reported (Table 3). The data show that Ba tends to dissolve mainly in calcium silicates (larnite, bredigite – from 0.5 up to 6 at.%) and in calcium aluminates (mayenite and brownmillerite – 0.5–40 at.%). V is mainly associated to brownmillerite (up to 1 at.%), larnite (approximately 1 at.%) and spinel (1.1 at.%), even significant amount can be detected also in wustite. Cr is mainly associated to spinel (30–50 at.%) and Ca-chromite (c.a. 50 at.%), and to a lesser extent to wustite (max 3 at.%) and brownmillerite (max 9 at.%). Traces of Cr can be also found in calcium silicates, especially larnite. In disagreement with the literature (Durink, 2008; Albertsson et al., 2013), Cr was detected neither in merwinite nor in akermanite-gehlenite.

Four main phases characterize almost all of the entire group of analyzed slag; the effects of such phases on leaching behaviour of fine particles will be discussed in the text. In particular, the influence of larnite, gehlenite and spinels is discussed. As documented by Belhadj et al. (2012) wustite is considered completely nonreactive, however.

1 Refer to Table 3 for abbreviation and formula.
Fig. 2 – Examples of microstructure: (a) sample #2; (b) sample #7; (c) sample #8 and (d) sample #12. Refer to Table 3 for abbreviations and formula.

Table 3 – Chemical composition, definition and toxic elements partition of the crystalline phases detected in the investigated slag samples (at.%).

<table>
<thead>
<tr>
<th>Phase</th>
<th>Symbol</th>
<th>Definition</th>
<th>Formula</th>
<th>Ba</th>
<th>V</th>
<th>Cr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bredigite</td>
<td>Br</td>
<td>di-calcium silicate</td>
<td>Ca$<em>2$(Mg$<em>x$Si$</em>{4-x}$O$</em>{10}$)$_4$ (7CaO·MgO·4SiO$_2$)</td>
<td>0.15–0.60</td>
<td>0.10–0.15</td>
<td>0.00–0.15</td>
</tr>
<tr>
<td>Brownmillerite</td>
<td>B</td>
<td>tetracalcium aluminoferrite</td>
<td>Ca$_2$(Al$<em>x$Fe$</em>{2-x}$)O$_3$ (4CaO·Al$_2$O$_3$·Fe$_2$O$_3$)</td>
<td>0.15–0.40</td>
<td>0.5–1.0</td>
<td>1.0–10.0</td>
</tr>
<tr>
<td>Ca–chromite</td>
<td>Ca–Cr</td>
<td>calcium chromite</td>
<td>CaCr$_2$O$_4$ (CaO·Cr$_2$O$_3$)</td>
<td>–</td>
<td>–</td>
<td>45.0–55.0</td>
</tr>
<tr>
<td>Gehlenite-akermanite</td>
<td>G</td>
<td>Ca–Mg silicate</td>
<td>Ca$<em>2$(Al$<em>x$Si$</em>{y}$)O$</em>{z}$–Ca$_2$MgSi$_2$O$_7$ (2CaO·Al$_2$O$_3$·SiO$_2$–2CaO·MgO·2SiO$_2$)</td>
<td>0.07–0.15</td>
<td>0.08–0.15</td>
<td>0.00–0.04</td>
</tr>
<tr>
<td>Kirschsteinite</td>
<td>K</td>
<td>calcium olivine</td>
<td>CaFeSiO$_4$ (CaO·FeO·SiO$_2$)</td>
<td>0.02–0.06</td>
<td>0.05–0.30</td>
<td>0.03–0.50</td>
</tr>
<tr>
<td>Larnite</td>
<td>L</td>
<td>di-calcium silicate</td>
<td>Ca$_2$SiO$_4$ (2CaO·SiO$_2$)</td>
<td>0.05–0.30</td>
<td>0.10–0.90</td>
<td>0.2–1.3</td>
</tr>
<tr>
<td>Mayenite</td>
<td>CA</td>
<td>calcium aluminate</td>
<td>Ca$_2$Al$<em>2$O$</em>{12}$ (12CaO·7Al$_2$O$_3$)</td>
<td>0.05–0.40</td>
<td>0.01–0.20</td>
<td>0.1–0.10</td>
</tr>
<tr>
<td>Merwinite</td>
<td>Mw</td>
<td>Mg–Mg silicate</td>
<td>Ca$_2$MgSi$_2$O$_7$ (3CaO·MgO·2SiO$_2$)</td>
<td>0.10–0.15</td>
<td>0.05–0.10</td>
<td>–</td>
</tr>
<tr>
<td>Mg–Cr–spinel</td>
<td>Sp</td>
<td>magnesiochromite</td>
<td>MgCr$_2$O$_4$/ (MgFe)(Al,Cr)$_2$O$_4$ (Fe,Mg,Fe)O/MgO·FeO</td>
<td>0.05–0.10</td>
<td>0.10–1.10</td>
<td>31.0–55.0</td>
</tr>
<tr>
<td>Wustite</td>
<td>W</td>
<td>iron oxide</td>
<td></td>
<td>0.02–0.10</td>
<td>0.05–0.70</td>
<td>0.5–3.0</td>
</tr>
</tbody>
</table>

Fig. 3. As expected, the high surface–on-volume ratio of the fine slag particles induced a higher release of Ba and V than that obtained performing the bulk test. In fact, with respect to the bulk slag, leached Ba was 3 times higher (Fig. 3a) and leached V was about twice as high for the fine particle samples (Fig. 3b). On the contrary, Cr leaching showed an opposite trend, since the highest Cr concentrations were produced by samples with very low Cr leaching during the standard test on 4 mm particles (Fig. 3c). Contrary to what was expected, the leachate pH was almost the same for powder and bulk slag (Fig. 3d). Only for some samples the powder slag induced a pH 5–10% higher than the respective bulk sample. This result emphasizes that the high S/V ratio of the fine particles favours a greater release of Ba and V and a concomitant reduction of Cr if compared with the marginal effect of the reduction of size on the final leachate pH.

These results provide a preliminary indication about the risk associated to the presence of fine particles during the production of aggregates, since the leaching behaviour could change extremely by the volume fraction of such fine powders. Nevertheless, the mass loss registered in both cases was negligible, hardly overcoming the 1 wt.%. This again underlines the limited interaction between slag and water and the primary role of the distribution of the phases within the slag.
Leaching tests on the slag powder confirmed the Ba-V leaching behaviour described by the same authors (Mombelli et al., 2016). When barium is detected, vanadium concentration is always under the ICP-OES detectable limit (50 μg/l).

Different relations in leachate concentration can be highlighted: Ba is directly proportional with Ca and Al dissolved in the eluate; inverse logarithmic dependence was identified between leached Mg and Ba (Fig. 4a). Dissolved calcium could derive from the dissolution of larnite, or other hydraulic silicates, as well as from brownmillerite or other Ca-rich compounds (mayenite). A study conducted by Engström et al. (2013) on pure slag minerals, confirms the attitude of larnite, mayenite and calcium aluminate to dissolve rapidly at all the investigated pH. Aluminium could be associated to calcium-aluminates, brownmillerite or spinels dissolution. In particular, the concentration of the above mentioned elements are not stoichiometric with respect to their concentration in the associated phases, thus complicating the identification of the involved phases. In fact, during the leaching of silicate-like material, the incongruent dissolution phenomenon generally occurs. This phenomenon, which was widely documented in the natural weathering process of silicate minerals by Ruiz-Agudo et al. (2012), leads to the formation of a Ca-depleted solid layer during leaching (Huijgen et al., 2006), in which the composition influences the dissolution/precipitation mechanisms, indicating non-stoichiometric release of solid phase constituents into the liquid phase.

However, the barium release seems to be effectively ruled by larnite, brownmillerite and mayenite dissolution, since these three mineralogical phases have hydraulic properties and good affinity with water (Lea, 1970; Taylor, 1990; Hewlett, 1998; Bensted, 2002; Gartner et al., 2002; Raab et al., 2008).

During hydration, brownmillerite forms $4\text{CaO} \cdot \text{Al}_2\text{O}_3 \cdot \text{nH}_2\text{O}$ and hydrated iron oxide gel (Hewlett, 1998; Taylor, 1990). This is a fast and exothermic reaction, but precipitation of an insoluble layer of hydrated iron oxide on the calcium aluminaferrite crystal surface forms a barrier to further reactions. For this reason the role of brownmillerite could be only marginal, even if the direct proportion between Ca and Al (Fig. 4b) seems to confirm the contribution of brownmillerite and mayenite to the leaching process.

Mg and Ba belong to the same chemical group and that can explain the detected reciprocal behaviour. Ba leaching seems to be favoured when Mg is well bound in non-hydraulic phases (i.e. wustite), whereas Ba leaching is limited when Mg might be released, and this is due to its implication in hydration phenomena. In addition, the solubility of Mg influences the solubility of Ba. For example, when the Mg concentration is high, the media is no longer able to dissolve Ba. The same behaviour is presented between Ca and Mg (Fig. 4b).

Vanadium leaching was associated to Cr leaching by linear relation, highlighting the mutual controlling behaviour of such species. Mg-V presents the same behaviour shown by Mg-Ba (Fig. 4c).

High leaching of Al, Ba, Ca, Cr and V was associated to high leachate pH, as indicated in Fig. 5a–c, whereas high Mg leaching were found in less caustic solutions (Fig. 5a). High concentrations of alkaline elements rapidly rise up the pH of a solution. In addition, high concentrations of Ca and Ba in the leachate could inhibit the dissolution of Mg, since its solubility is inversely proportional to other alkali concentrations. V and Cr behave as anions, and should be highly dissolved in basic pH, as reported in U-Mate report (U-Mate, 2012). The leachate conductivity shows the same trend of pH as a function of the main elements investigated (Al, Ba, Ca and Mg), since high
Fig. 4 – Interaction among leached substances: Ba vs Mg, Ca and Al (a); Ca vs Al and Mg (b); V vs Mg and Cr (c).

Fig. 5 – Effect of leached substances on leachate pH: Ba and Mg (a); Ca and Al (b); V and Cr (c).

Fig. 6 – Effect of leached substances on leachate conductivity: Ba and Mg (a); Ca and Al (b).
ions concentration in a solution lead to an increase of electrical conductivity (Fig. 6a and b). No correlation amongst Cr, V and conductivity were found.

3.3. Effects of microstructure on leaching in standard conditions

Correlation between elution test results and crystallographic analysis allows to better determine the potential phases responsible for the elements being released.

A logarithmic trend was identified between larnite fraction and the leachate pH (Fig. 7a). This relation seems to validate the hypothesis about the key role played by larnite in slag leaching behaviour: larnite fraction controls the most of Ca concentration in the leachate (Fig. 7b) and it also determines the final water pH. The higher the larnite fraction in the slag, the higher the ion concentration (Al, Ca and Ba) in the leachate. This is confirmed by the direct proportion between larnite fraction and leachate conductivity (Fig. 7a). Even if Al is not an element-forming of larnite, the occurrence of incongruent dissolution can explain why Al leaching is high in high larnite fraction slag.

Mg dissolution manifests an opposite trend as the larnite fraction increases. This result confirms the role of larnite in Ba leaching and the observed trend can be explained on the basis of the hydration chemistry of dicalcium silicate, discussed in the previous paragraph.

Slag characterized by a high gehlenite fraction signal a significant reduction in calcium, aluminium and barium elution (Fig. 7d). High gehlenite fraction could induce modestly high Mg leaching and prevent the Ba leaching. The dissolution of BaO into the gehlenite reduces the solubility of MgO in the same phase. In addition, the amount of Ba and V trapped in gehlenite (Table 3) is not leached, since this phase has been proved to own extremely low solubility in water solution, especially at high pH (Engström et al., 2013). If the slag is poor in chemical species that can bind MgO, Mg leaching can be enhanced. In fact, high Mg elution is associated to low Mg-Cr-spinels fraction (Fig. 8a), as well as AI leaching.

The spinel fraction seems to control the leaching of V and Cr (Fig. 8b). For instance, the higher the spinel fraction, the higher the V and Cr leaching is. High spinel fraction is generally associated to the slag belonging to group B, hence characterized by high Cr2O3 content (5-30 wt.%). In fact, as reported by Liao et al. (2016), the Cr incorporation in the spinel phase depends upon its initial concentration in the slag. Higher the available Cr, higher the amount of Cr incorporated in the spinel is. Moreover, the B-group slag are generally characterized by a low amount of MgO and Al2O3 (Table 2) and thus, the spinels tend to be over-saturated in Cr. Attempting to evaluate the spinel stability of such slag thorough the use of sp-factor (Eq. (1)) (Kühn et al., 2006; Mudersbach et al., 2009), the excess of Cr2O3 and the low amount of MgO and Al2O3 play against the formation of stable spinels.

$$sp\text{-factor} = 0.2 \cdot \text{MgO} + 1.0 \cdot \text{Al}_2\text{O}_3 + n \cdot \text{FeO}_n - 0.5 \cdot \text{Cr}_2\text{O}_3 \quad (\text{wt.\%})$$

where n is the optical electronegativity of iron ions, depends on the oxidation state of the EAF-slag and can assume a value of between 1 and 4 as a function of the oxidation number assumed by the Fe ions.

Furthermore, it seems that the higher the spinel fraction, the higher the probability is to form unstable spinel. This hypothesis seems confirmed by the analysis reported in Fig. 9, related to three different samples, one belonging to group A
Fig. 8 – Effect of spinel weight fraction on Mg, Al (a) and V, Cr (b) leaching.

Fig. 9 – Comparison between the spinel formed in three different slags belonging to (a) group A (sample A) and (b, c) group B (samples B1 and B2).

| [atomic %] Spinel chemical composition (SEM-EDS) |
| Ca | Si | Al | Mg | Cr | Fe | V |
| 0.64 | 23.45 | 11.47 | 41.93 | 21.98 | 0.93 |

Standard leaching test
Cr: 0.104 mg/l; V: 0.963 mg/l

Standard leaching test on powder
Cr: 0.019 mg/l; V: 0.12 mg/l

Spinel fraction: 5-10\%wt.

| [atomic %] Average chemical composition (ED-XRF) |
| CaO | SiO\(_2\) | Al\(_2\)O\(_3\) | MgO | Cr\(_2\)O\(_3\) | FeO | V\(_2\)O\(_5\) |
| 16.61 | 21.42 | 10.27 | 2.3 | 3.48 | 17.76 | 0.16 |

As reported by Albertsson et al. (2013) and Arredondo-Torres et al. (2006), a slag with high basicity is interested by spinel solid solution ((Mg,Ca)Cr\(_2\)O\(_3\)) precipitation from the matrix with high amount of impurities and irregular structure and morphology. This seems attributable by the presence of Cr\(_{6+}\) that induces spinel instability. Cr\(_{6+}\) formation is favoured in high basicity slag, since they should have high fraction of not combined CaO that could react with Cr to yield Cr\(_{6+}\) (Lee and Nassaralla, 1998). Spinel instability due to high basicity (and thus high Cr\(_{6+}/Cr^{3+}\) ratio) has been proved to induce high Cr leaching during standard leaching test (Mombelli et al., 2016). In addition, the low Al\(_2\)O\(_3\) content in the spinel analyzed in the sample B1 and B2 contributes to reduce its stability, thus leading to significant Cr leaches. These aspects have two consequences: even if the chromium is mainly bound in spinel form, the chromium leaching still occurs, and then, the excess of Cr decreases the solubility of V into the spinel phase, favouring its mobility and thus the leachability of such an element (Fig. 9).

### 3.4. Leaching test at different L/S ratio

The set of elution tests indicated that at a standard L/S = 10 l/kg the interaction between water and slag is minimal and the releasing behaviour is mainly governed by the high surface-to-volume ratio of the powder. The weight loss was very low and only in few cases overcome 1% even if the high powder surface-on-volume ratio should enhance the kinetic reactions and dissolution rate. This result implies that only a few portions of the material interacts with water and thus, the identification of the dissolved phases during the leaching test (by XRD or SEM) is difficult. Only by increasing the liquid-on-solid ratio is the dissolution of a greater amount of slag possible. Nevertheless, slag dissolution is strictly related to microstructural features and crystallographic phase...
distribution, since not all the structural constituents have the same reactivity with water (Engström et al., 2013; Strandkvist et al., 2015).

The effects of the liquid/solid ratio were quite evident for all of the investigated samples: increasing the water volume caused a corresponding increase in the amount of dissolved slag (Fig. 10a). By increasing the L/S ratio, the leachate pH decreased (Fig. 10b), as well as the leachate conductivity (Fig. 10c). With the water being less caustic at a high liquid-to-solid ratio, compared to the standard condition, the release of dangerous chemical species could be potentially heightened as the environmental conditions remain away from saturation limits (Mombelli et al., 2014; Nicoleau et al., 2013).

To identify which phases are involved in the hydration/dissolution process, and thus leading to the pollutants release, elution tests on powdered samples at different liquid-on-solid ratio were carried out. As described in Fig. 10a, only at a high L/S ratio slag mass is significantly dissolved. For this reason, at 10 l/kg, it is quite impossible to detect any variations in the XRD pattern. Fig. 10 shows the results of XRD analysis performed after the elution test at different L/S on several slag samples. In order to compare the different spectra, normalization on the more intense wustite peak is performed. Wustite is selected because it should not react with water; in other words its dissolution rate is so low that it can be considered insoluble (Belhadj et al., 2012). The XRD patterns in Fig. 11 demonstrate the phases involved in the leaching process, confirming the results discussed in the previous paragraphs. The different XRD spectrums resolve that, using a standard L/S ratio (10 l/kg), the interaction between slag and water is only cortical and interests a limited portion of material. Larnite and brownmillerite peaks decreased their intensity at high L/S ratios (Fig. 11b and d). In addition, Fig. 11b showed the alteration of Ca-chromite, demonstrating the implication of such a phase in the Cr leaching, previously discussed by Mombelli et al. (2016) on the basis of slag basicity. This structural constituent probably contributes to the uncommon chromium concentration in the leachate, because the reaction between CaO and Cr oxide fixes the Cr status to hexavalent form. In alkaline circumstances and in the presence of atmospheric oxygen CaCr2O4 can easily be oxidized to CaCrO4, a compound that easily leaches chromium and builds hexavalent chromium from divalent or trivalent chromium (Meegoda et al., 2008; Schwandt and Fray, 2006; Ylipekkala, 2005). The high value of CaCrO4 solubility constant indicates the high solubility of this compound (Selected Solubility, 1997). Also the spinel peaks did undergo intensity reduction, even if the variation is less evident. However, the same analysis highlighted the stability of gehlenite and kirschsteinite, even at a high liquid-to-solid ratio. Thus, these phases can be classified as preferred crystalline phases, able to ensure high microstructural stability (Fig. 10a and d). These results comply and validate those presented by the same author (Mombelli et al., 2014).

Among the different elution test trials, only weak intensity peaks related to the formation of typical hydrated phases

| Table 4 – SEM-EDS chemical composition of the phases indicated in Fig. 12 (at.%). |
|-----------------------------------|---|---|---|---|---|---|---|---|---|---|---|---|
| at.%                          | Mg | Al | Si | P | S | Ca | Ti | V | Cr | Mn | Fe | Ba |
| Before leaching test          |    |    |    |   |   |    |    |   |    |    |    |    |
| General                        | 7.36 | 6.78 | 18.81 | 0.11 | 0.15 | 17.92 | 0.29 | 0.11 | 3.98 | 5.45 | 38.98 | 0.06 |
| W                              | 7.94 | 6.78 | 18.81 | 0.11 | 0.15 | 17.92 | 0.29 | 0.11 | 3.98 | 5.45 | 38.98 | 0.06 |
| K                              | 10.50 | 40.69 | 35.66 | 0.21 | 41.83 | 5.01 | 27.31 | 0.11 |
| Sp                             | 8.84 | 16.29 | 0.40 | 0.21 | 41.83 | 5.01 | 27.31 | 0.11 |
| G                              | 3.40 | 23.10 | 29.95 | 38.78 | 0.72 | 9.96 | 0.02 |
| After leaching test            |    |    |    |   |   |    |    |   |    |    |    |    |
| General                        | 8.09 | 7.67 | 18.52 | 0.17 | 0.30 | 18.01 | 0.15 | 3.98 | 4.86 | 38.14 | 0.11 |
| W                              | 10.28 | 7.67 | 18.52 | 0.17 | 0.30 | 18.01 | 0.15 | 3.98 | 4.86 | 38.14 | 0.11 |
| K                              | 12.70 | 43.59 | 33.62 | 2.42 | 7.67 | 0.72 | 4.05 |
| Sp                             | 11.03 | 24.50 | 2.34 | 33.00 | 4.01 | 25.12 |
| G                              | 4.32 | 23.99 | 28.86 | 31.33 | 1.52 | 9.96 | 0.02 |
Only for the C group samples the high free lime fraction allows a clear identification of portlandite peaks after the leaching test performed on the slag powder. In order to confirm the non-hydraulic behaviour of the above mentioned phases (gehlenite, kirschsteinite), an elution test with extreme L/S ratios was performed on several samples characterized by gehlenite-kirschsteinite matrix. SEM-BSE microographies did not show any differences before and after the test (Fig. 12). The slag structure appears unaltered and the chemical composition of the phases measured by the EDS probe showed a negligible calcium and iron dissolution (Table 4).

These results highlight the important role of crystalline phases in slag leaching, as previously stated by several authors (Chaurand et al., 2007; van Zomeren et al., 2011; Manso et al., 2005; Geiseler, 1996; Mombelli et al., 2014; Gelfi et al., 2010; Engström et al., 2014; Baciocchi et al., 2015; Costa et al., 2015). While the microstructure is strictly related to the slag chemical composition, a prediction on the leaching attitude of the slag can be evaluated by coupling chemical information, CaO-SiO$_2$-FeO$_x$ and CaO-SiO$_2$-Al$_2$O$_3$ ternary diagrams. In fact, ternary diagrams provide reliable indications about the crystalline phase/phases in the slag matrix, as established by Mombelli et al. (2016). The dangerous metals involved in the leaching process from EAF carbon steel slag are controlled
by specific crystalline phases (i.e. larnite, brownmillerite, etc.) formed in the slag as a consequence of its chemical composition and cooling method.

4. Conclusions

In this work the effects of crystalline phases on slag leaching behaviour were investigated, allowing for the definition of the leaching mechanisms involved in the release of polluting elements. The analysis of fine particles allows for a clarification of the effect of fine fraction on leaching and to identify what phases are involved in the dissolution process. Liquid-on-solid ratio plays a key role on slag leaching behaviour: an increase in L/S can lead to higher slag dissolution and a higher release of pollutants than in standard conditions. Thus, the characterization of the slag leaching behaviour through standard leaching tests alone could be misleading in order to assess the safety of this by-product. If, in fact, if slag is used as unbound stone material, the continuous wetting-drying cycles may lead to continuous dangerous substances leaching due to layer-by-layer dissolution. This consideration suggests characterize slag not only by standard elution test but also with a modified procedure considering the effect of liquid-on-solid ratio. Similarly, the fine particle content can significantly modify the environmental conditions, increasing the leaching rate, even if the slag were declared safe after standard tests. This suggests setting-up leaching tests with different particle sizes.

Mineralogical phases in the slag microstructure strongly influence the slag leaching. Larnite (and its polymorphism) are the phases mainly dissolved during elution tests (in standard conditions) and are responsible for Ba and V releasing, also controlling the leachate properties (pH and conductivity). Brownmillerite has less solubility than larnite but contributes to an increase in Ba and Cr content in eluted water. Chromite-like phases, and not stoichiometric Mg-Cr-spinel, control the chromium dissolution.

High stable phases, like gehlenite and kirschsteinite, seem to improve the slag resistance against dissolution, also at high L/S ratio, and these are thus the structural constituents to be favoured in order to produce safe and environmentally friendly slag. Even with different characteristics, in all the investigated slag groups, high gehlenite fraction is associated with better behaviour against pollutant leaching, producing a less caustic and conductive leachate. Thus, gehlenite development is the goal to be achieved in order to enhance the recycling of carbon steel slag as artificial aggregates for bound and unbound applications. The industrial processes developed to stabilize/inertize the EAF slag confirmed this statement. In fact, they are designed to foster the formation of a gehlenite matrix after slag cooling by adding silica or other additives to the molten slag (Patent, 2012, 2014).

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References


