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The effect of chemical composition on the leaching behaviour of electric arc furnace (EAF) carbon steel slag during a standard leaching test

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ABSTRACT

The electric arc furnace (EAF) slag could be exploited in several fields of application, such as land filling, road constructions and concrete production. However, their use is limited by the presence of polluting chemical elements (chromium (Cr), barium (Ba), Vanadium(V), etc.) that can be dangerous to humans and the environment. Thus, chemical and structural stability is a fundamental requirement, especially when the slag may come in contact with water. Therefore, the interaction between slag and water is key, in order to classify the slag as a safe raw material. In this work, the effect of slag chemical composition on the chemical leaching of about seventy EAF carbon steel slags of different production steel grades was investigated. Standard leaching tests (24 h at 10 l/kg) in deionized water on slag bulks were performed and the results were correlated with the slag chemical composition. The survey has made possible defining the safe chemical composition areas on the main ternary diagrams, able to transfer stability to the slag. The results obtained indicate the water/slag ratio as the most important factor in the release of polluting substances, also identifying a critical scenario for slag recycling.

Keywords:

Electric arc furnace (EAF) slag
Leaching behaviour
Standard leaching test
Chemical composition
Ternary diagrams

Introduction

Steel slags are today largely utilized in roads, as construction and paving materials, since their physical properties are similar, or sometimes better, than natural materials like gravel.

Environmental concerns regarding utilization and landfilling of steel slags focused on the content of heavy metals and particularly on the leachable quantities, where chromium (Cr) has gained special attention due to its toxicity in the hexavalent state [1,2]. Other chemical species, i.e. barium (Ba), vanadium (V), molybdenum (Mo), require monitoring due to their high toxicity on humans and the environment [3,4]. The identification of leaching behaviour is mandatory for slag use in unbound applications, i.e. unpaved roads, armourstone or gabions, because in such conditions, slag is subject to a continuous cycle of wetting and drying. Several studies [5–9] have investigated the leaching behaviour of slag aggregates, and highlight that such aggregates could potentially release dangerous chemical elements (especially Ba, V and Cr), as well as unbound slag, when coming into contact with water. In

particular, the recycling of such a slag is subordinated with respect to the severe limits imposed by national legislative decrees that change from country to country in European Union. Also the list of substances and chemicals to be monitored changes from country to country. For examples, Italian environmental legislation requires the monitoring of 22 parameters, namely pH, COD, As, Ba, Be, Cd, Co, Cr(tot), Cu, Hg, Mo, Ni, Pb, Sb, Se, V, Zn, Cl⁻, CN⁻, F⁻, NO₃⁻, SO₄²⁻, whereas in Germany only few parameters must be controlled (pH, EC, Cr, Mo, V and Fluorides). In France and Spain almost the same elements are restricted (particularly focused on heavy metals). The limits on Cd and Se in Spain are the most strictly (less than 1 µg/l) amongst the considered European countries. Moreover, German legislation does not require any control on Ba leaching, whereas the Ba limits in France and Spain are twice the level of Italian limits. Also the V is considered differently in the various EU countries: severe limits are set in Italy, Germany and Spain, but not in France [10–15].

To determine the concentration of leached elements every country adopts a different test criteria that contributes to confusion about slag environmental safety (Table 1), in spite of the European Committee of Standardization (CEN) is considering the introduction of common harmonized standards [16,17].

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Table 1
Environmental legislation and leaching test standards for different European countries.

Country	Italy	Germany		France	Spain (Basque Country)
Environmental legislation	DLGS 152-2006	EBV	TL Gestein StB 04	SETRA Guide de laitiers sidérurgique EN12457-4	RD 34/2003 RD 49/2009 EN12457-3
Leaching test standard	EN12457-2	DIN19528 DIN19529	DIN38414-4		
L/S ratio	10 l/kg	2 l/kg	10 l/kg	10 l/kg	First step 2 l/kg Second step 8 l/kg
Sample granulometry	<4 mm	<32 mm	<32 mm or 8–11 mm	<10 mm	<4 mm
Test duration	24 h	24 h	24 h	24 h	First step 6 h Second step 18 h

In this study, the Italian regulation was taken as reference since very critical test criteria are adopted, with high liquid-to-solid ratio (L/S) coupled with a small aggregate size that could enhance the leaching of such a chemicals.

Even if the conditions under which the leachates are produced vary as a result of different physical parameters like pH, temperature, or flow conditions, and influence the resulting concentrations of metals in the leachates [18], the slag chemical composition covers an important role on immobilization, or the release of the above mentioned toxic elements. For example, MgO concentration in the slag seems to be favourable for Cr immobilization but detrimental for the leaching of other polluting elements, i.e. Ba and V [19]. The solution behaviour of these elements have a parabolic law as a function of CaO and MgO concentration, whereas their dependence on silica concentration is generally described by a linear law [20]. Currently, the role of chemical composition on carbon steel slag leaching behaviour has not been extensively and systematically investigated, and is often only dedicated to Cr leaching [21,22]. For these reasons, an in-depth understanding of the role of the average chemical composition on slag leaching behaviour in standard conditions (deionized water at pH 7, room temperature, L/S: 10 l/kg) could give important indications to steelmakers about how to properly manage or modify their slag to improve their environmental sustainability and recycling.

In this study, different classes of carbon steel EAF slag from different steel productions (reinforced bar steel, high alloyed steel and quality steel) were investigated, in order to correlate their leaching behaviour in standard conditions with chemical features. The main goal of the present work is to forecast the leaching behaviour of a slag by average chemical composition, and define a corrective path to transform slag into a safe by-product. Chemical composition and slag-water interaction were investigated and correlated with the results of standard leaching tests (in accordance with EN 12,457-2) performed on bulk slag 4 mm in size. The effects of the main chemical species (CaO, MgO, Al₂O₃, . . .) on Ba, Cr and V leaching is discussed. To better understand how the slag interacts with water during standard leaching tests, qualitative tests were performed on 4 mm of crushed slag grains at different L/S ratios (10 and 100 l/kg). The analyses allowed for the identification of safe chemical composition areas on the main ternary diagrams that ensure full slag stability.

Table 2
Slag classification and number of samples according to chemical composition and steel grades.

Group ID	No. of samples	FeO _x	Cr ₂ O ₃	Production
A	13	>25%	<5%	Reinforcing bar steel
B	10	>25%	>5%	High alloyed steel
C	46	<25%	<5%	Quality steel

Experimental procedure

The EAF slags were provided by different Italian and European electric steelworks and associated with different steel productions. Sixty-nine different samples were collected and investigated, classifying the different slag into three groups according to the amount of iron and chromium oxide (Table 2). The number of samples for each group is indicated in Table 2.

The slag chemical composition was determined by ED-XRF analysis through the Ametec Spectro Xepos spectrometer in He atmosphere on 5 g of powdered slag.

The average chemical composition of the slag is reported in Table 3, and their position on ternary diagrams is shown in Fig. 1.

A morphological and microstructural investigation was performed by means of XRD and SEM analyses.

X-ray diffraction (XRD) data were collected using a Bruker D8 Advance diffractometer in a θ - θ configuration employing the Cu K α radiation ($\lambda = 1.54 \text{ \AA}$) with a fixed divergence slit size 0.5° and a rotating sample stage. Grinded samples (average diameter $15 \mu\text{m}$) were scanned between 10 and 80° (step size of 0.007°) with the Vantec detector.

The morphological and microstructural characterization was performed by Zeiss EVO50 scanning electron microscopy (SEM) equipped with an Oxford Inca EDS probe. The slag was moulded in araldite-based resin, before being ground and polished.

Standard leaching tests were performed according to EN 12,457-2 on granulated slag ranged from 4 to 0.1 mm. The fine fraction below 0.1 mm was discharged. The slag test portion was brought in contact with fixed volume of deionized water (10 l/kg) and stirred at 10 rpm by a rotatory mixer for 24 h. The test is based on the assumption that equilibrium or near-equilibrium is achieved between the liquid and solid phases during the test. The concentration limits taken as a reference are indicated in the Italian legislative decrees (D.M. 03 August 2005 N. 201 [10] and D. M. 05 April 2006 N. 186 [11]). Leached concentrations were measured by ICP-OES (induced coupled plasma-optical emission spectroscopy, detection limit $30 \mu\text{g/l}$).

To investigate slag-water interaction, a crushed particle (approximately 4 mm in size) for every slag was leached as a standard leaching test required but varying the L/S ratio (10, 100 l/kg). Slags were characterized by SEM, before and after elution tests, to detect morphological alterations, and to make a qualitative comparison on the effects caused by water erosion. The same

Table 3
Chemical composition range (min–max) of the investigated slag (% weight).

Group ID	MgO	Al ₂ O ₃	SiO ₂	CaO	FeO _x	Cr ₂ O ₃	V ₂ O ₅	Ba
A	2–5	10–15	15–20	15–25	30–50	2–5	0.1–0.2	0.05–0.1
B	1–3	1–3	5–25	15–25	30–50	5–30	1–2	ND
C	5–15	5–15	10–40	20–50	5–30	0.5–5	0.05–0.4	0.1–0.5

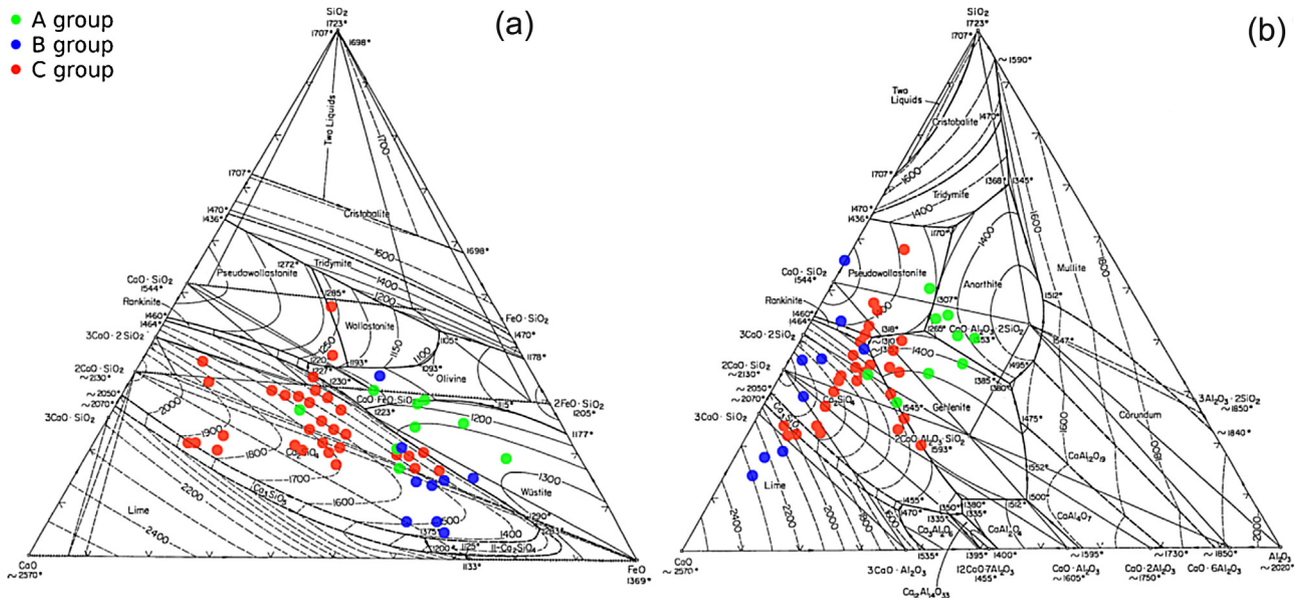


Fig. 1. Sample identification on (a) CaO–SiO₂–FeO and (b) CaO–Al₂O₃–SiO₂ phase diagram [23]. Some points overlap.

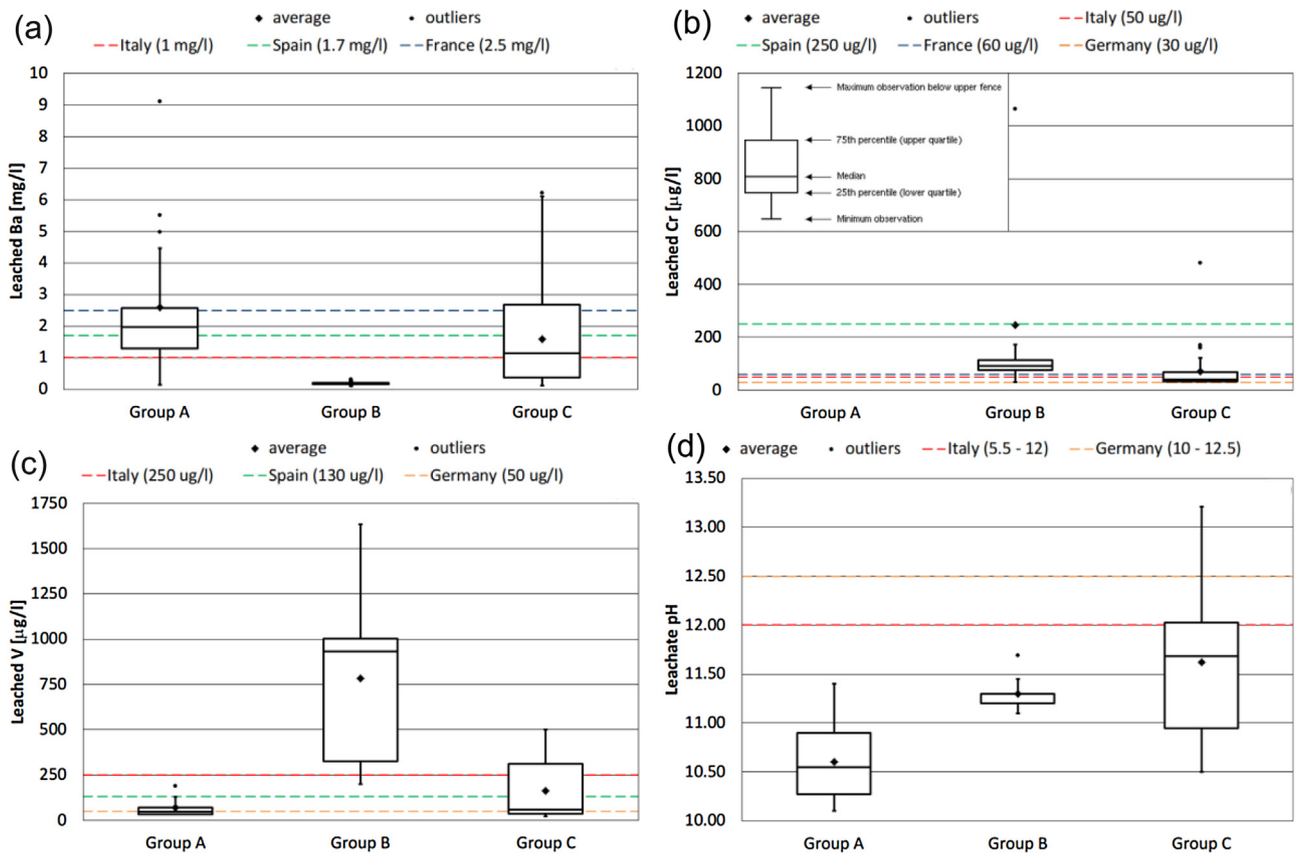


Fig. 2. Leaching data boxplots of investigated EAF carbon steel slag: (a) Ba, (b) V, (c) Cr and (d) pH. The concentration limits indicated are the most strict provided by the different environmental laws applied in the different countries.

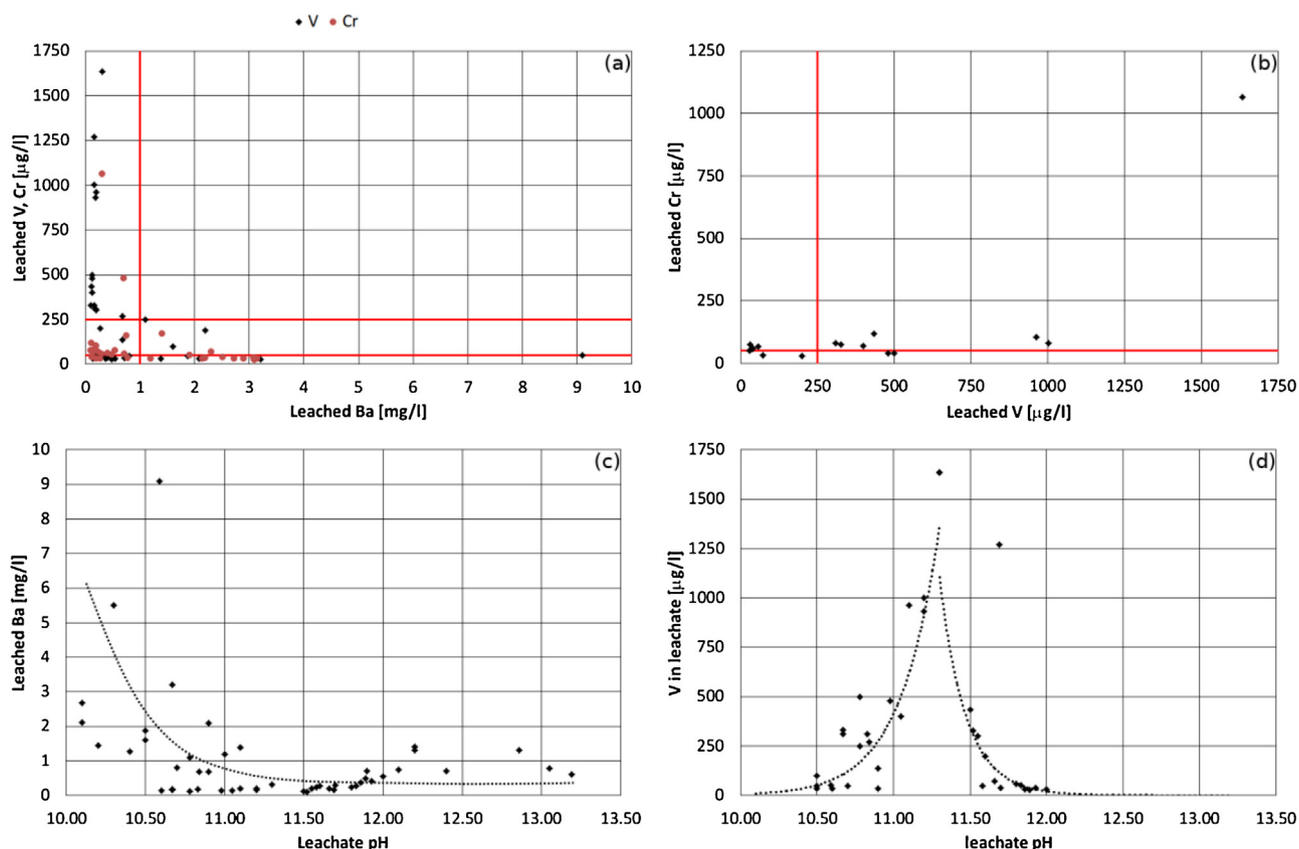


Fig. 3. Comparison amongst Ba, V, Cr and pH concentrations in leachate: (a) Ba–V and Ba–Cr relationships; (b) V–Cr relationship; (c) pH–Ba and (d) pH–V relationships. Red lines refer to Ba, V and Cr Italian leaching limits, 1 mg/l, 250 µg/l and 50 µg/l respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

particle was repeatedly treated at the different L/S ratio and the same area was analysed after each condition.

Results and discussion

Chemical, crystallographic and microstructural characterizations

Other differences, with respect to the first classification, can be highlighted amongst the different slag typologies. In fact, slag from the reinforcing bar steel production (group A) is characterized by high iron (averagely 40% wt.) and aluminium oxide (above 10% wt.) content (Table 3). High oxygen flow is required during steel production due to the high heterogeneity of the scrap used, in terms of size and chemical composition and the high content of polluting elements.

The slags belonging to group B are generally typified by a low content of MgO and Al₂O₃ (less than 3% wt.), and by a high fraction of Cr, V and Mo oxides, derived from both the high alloyed scrap used during manufacturing and from the ferro-alloys additions during the refining (Table 3).

The slag associated to quality steels (group C) is mainly formed by CaO (sometimes over the 50% wt.) and a small quantity of FeO (limited oxidation of the steel scraps to limit the FeO around 15–20% wt.) (Table 3). Quality steels require a very low P and S content, and therefore a large amount of CaO is necessary to maintain high slag activity.

Despite of the significant differences in chemical composition, crystallographic and microstructural analysis of the investigated

slag highlighted the occurrence of five main phases¹ [24–36]: wustite, (Mg, Cr, Fe, Al)-spinel, bredigite and/or larnite, brownmillerite, gehlenite.

Other phases, like kirschsteinite, were detected, generally in a sample featuring high FeO content and a moderate MgO fraction (group A and B), whereas merwinite is formed in slag with high levels of MgO and CaO (group C). Slags belonging to group C were also characterized by a small fraction of tri-calcium silicate and free lime, since these samples are characterized by the highest CaO content that cannot be completely complexed by SiO₂. In two samples (one belonging to group B and one to group C) the XRD pattern pointed out a particularly intense peak associated to Ca-chromite.

The role played by these crystalline phases during leaching tests is discussed in detail in the following paragraphs.

Standard leaching test results

Standard leaching test results, performed according to EN 12,457-2, on slag particles ranged from 4 to 0.1 mm highlight that carbon steel EAF slag manifest only Ba, V and Cr leaching, whereas the other species were always under the regulatory limits [37]. These results were observed also in previous experiments [38–40] and for this reason only Ba, Cr and V behaviour were discussed. The Ba, V and Cr mean and median values surpassed not only the maximum Italian concentration permitted, but also, in some cases,

¹ Refer to “LIST OF PHASES” for the chemical formula.

those imposed by the other European countries [12–15] (Fig. 2). Group A slag only seemed to be problematic for Ba, whereas V and Cr values are always far enough from regulation limits (Fig. 2a). Group B was mainly characterized by V leaching even if some samples also induced Cr leaching (Fig. 2b). Group C slag showed Ba and Cr release, whereas V, in most of the samples, complied with the limit, even if all the vanadium content characterizing the steel scraps is absorbed by the slag in oxidized form (Fig. 2c). Even if V does not seem problematic for this last slag group, the detection of high V concentration released by some samples does not exclude V leaching for this kind of slag. This result indicates that Cr release is always associated to the release of other elements, namely V, whereas Ba leaching can occur with or without the leaching of other substances (Fig. 3). Moreover, Ba and Cr leaching seems to interest carbon steel slag the most (A and C groups), and V leaching is only prevalent in high alloyed steel slag (B group).

Some outliers were identified for Ba and Cr. In the first case, three samples of group A released an unexpected high concentration of Ba. Currently, no explanations have been found, since the microstructure of those samples did not indicate particular phases or unusual configurations. This behaviour could depend on the high MgO content and will be discussed in the next paragraphs. The outlier samples that leached an unexpected concentration of Cr belong to group B and group C, respectively. The samples are those featured by Ca-chromite and this phase is probably the main phase involved in Cr leaching.

The leachate pH complied with the safety range proposed by the regulation in most of the analysed samples, and its average values increase in a linear form from group A to group C, and is probably due to the different content of CaO and FeO in the slag (Fig. 2d). Leachate from group C reached the highest pH values and in some cases surpassed the maximum admitted value (12 pH). Over that value the environment becomes caustic and must be classified as corrosive, further limiting the possibility to use steel slag for civil purposes [41].

Ba concentration was high in leachates featured by low pH numbers. Ba in the slag is mainly present in the form of BaO and, since it is an alkaline oxide, it tends to dissolve first in water together with the other alkaline elements. However, if the environment reaches alkaline pH, the BaO migration should be reduced [42]. This hypothesis is consistent with the experimental evidence (Fig. 3c) and should explain the low Ba concentrations when the leachate reached high pH values (between 11 and 13).

Vanadium oxide is considered an amphoteric species [43,44] and in alkaline solutions tends to behave as acid oxide [45]. Since the EAF slag is basic slag, the leachate produced from their dissolution generally reach a pH in the range 11–13. This could

explain the high concentration of such elements in leachate having a pH in the range 11–11.5 [46]. However, when the pH reaches very high values (above 12), the solubility of V can be drastically reduced (Fig. 3d).

Ba and V seem to have an opposite tendency when dissolved in water (Fig. 3a). Firstly, no samples manifested Ba and V concentration over the imposed limits in the leachate at the same time. In fact, in all samples with Ba leaching, V complies with the limits, and vice versa. This behaviour could depend on, and contemporary control, the final leachate pH and should be ruled by slag chemical composition and phase distribution. In the graph in Fig. 3, it is possible to note some slag fall in the “safe area” (confined between V and Ba regulation limits). Those slags are generally characterized by a crystalline gehlenite microstructure that traps and inhibits chemical leaching, due to poor hydraulic properties [47–50].

These results suggest that the correlation between chemical composition and leaching test results could help identify some common aspects governing the interaction between slag and water. For example, the correlation between Cr₂O₃ content in slag and Cr concentration in the leachate can be excluded. In fact, B and C slag leachates had almost the same average Cr concentration even if the Cr₂O₃ content of the first group is five times more than the latter ones. In addition, the same conclusion can be reached for iron oxide, since different FeO/Fe₂O₃ content featured in the different slag groups.

Leaching test on crushed slag particles

Different slag belonging to groups A, B and C were investigated, and some examples are reported in Figs. 4 and 5. The same particle was repeatedly treated at the different L/S ratio and the same surface was analysed after each test condition. The liquid-to-solid ratio is one of the fundamental factors ruling the interaction between slag and the environment. Its effect is evident: increasing the water volume caused an increase in the amount of dissolved slag (Fig. 4). In fact, in standard conditions, the interaction between water and slag was superficial and is only of interest to a limited portion of the surface particle (Fig. 4b), but by increasing the L/S ratio, the surface of the slag particle appeared to be largely consumed (Fig. 4c). This implies that during the standard test, mass loss is not appreciable, even if the slag-water interaction is enough to change the leachate pH and drive heavy metal leaching. However, not all the surface reacts with water, but only specific phases that are involved in hydration and dissolution processes. In particular, SEM-EDS analysis on the surface (before and after the test) indicated that the Ca-rich silicates, i.e. larnite and

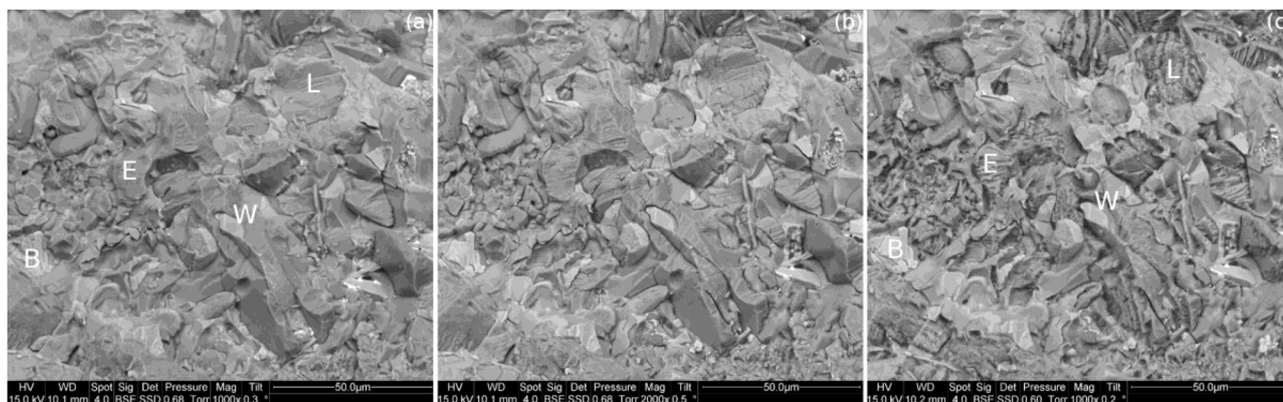


Fig. 4. SEM-BSE micrographies of crushed slag particles (belonging to group C) leached at different L/S ratios: (a) as-received; (b) 10 l/kg; (c) 100 l/kg. B: brownmillerite; E: eutectic 3CaO·SiO₂-FeO; L: larnite and W: wustite.

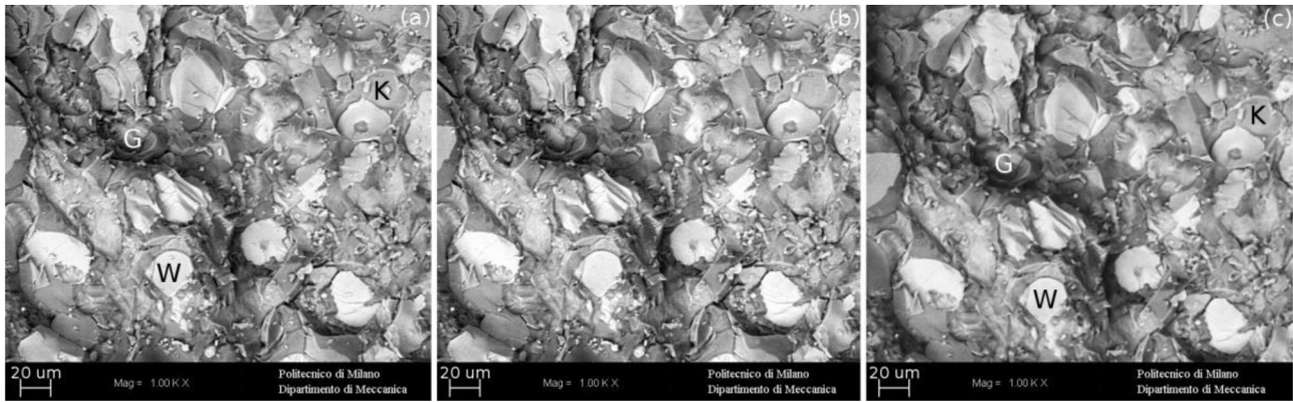


Fig. 5. SEM-BSE microographies of crushed slag particles (belonging to group A) leached at different L/S ratios: (a) as-received; (b) 10l/kg; (c) 100l/kg. G: Gehlenite; K: kirschsteinite; W: wustite.

Table 4

SEM-EDS point analysis of crushed slag particles before and after leaching test at 100l/kg (% atomic). B: Brownmillerite; E: eutectic $3\text{CaO}\cdot\text{SiO}_2\text{-FeO}$; L: larnite; G: gehlenite; K: kirschsteinite; W: wustite.

Sample	Phase	Condition	Mg	Al	Si	Ca	Ti	V	Cr	Mn	Fe	Ba	
Fig. 4	W	Before leaching	14.27			19.83			1.87	20.45	43.48	0.11	
		After leaching 100l/kg	16.30			18.02	0.02		2.11	20.15	43.39	0.01	
	E	Before leaching		2.25	22.96	66.95		0.05	0.67	2.45	4.48	0.20	
		After leaching 100l/kg	4.04	5.99	14.93	28.59	0.44	0.14	2.71	15.22	27.85	0.08	
	B	Before leaching	1.00	25.79	1.18	49.09	1.41	0.43	8.48	1.03	11.47	0.10	
		After leaching 100l/kg	1.03	23.28	2.46	48.45	1.60	0.36	10.50	1.53	10.79		
L	Before leaching		1.49	31.22	66.01		0.38	0.23		0.50	0.17		
	After leaching 100l/kg	3.49	4.66	4.37	29.26					5.02	53.20		
Fig. 5	W	Before leaching	8.34					0.06	1.00	9.06	81.52	0.02	
		After leaching 100l/kg	10.35			0.43		0.10	1.10	8.81	79.15	0.05	
	K	Before leaching	10.30		39.79	36.67					3.30	9.94	
		After leaching 100l/kg	12.50		43.69	33.72					2.32	7.70	0.07
	G	Before leaching	3.28	21.98	29.86	39.78					0.66	4.44	
		After leaching 100l/kg	4.25	23.08	28.53	31.34					1.40	11.38	0.01

brownmillerite, drive the slag dissolution (B and L marks in Fig. 4). The dissolved larnite fraction rapidly increases whereas brownmillerite dissolution is clearly evident only at high L/S ratios. Wustite (W) remains unaltered whereas in the complex eutectic phase $3\text{CaO}\cdot\text{SiO}_2\text{-FeO}$ (E) a large amount of CaO and SiO_2 were dissolved, creating a characteristic striped morphology (Table 4). Similar results were obtained for the other slag samples, fully in agreement with those previously reported by other authors [51,52].

However, not all the investigated slag behaved in the same way. For example, the surface of the slag reported in Fig. 5 appeared completely unaltered after immersion at 100l/kg. This slag complied with leaching limits, thus enabling the possibility to discern between safe and unsafe slag. In this case, the slag microstructure was mainly formed by kirschsteinite (K mark in Fig. 5), gehlenite (G) and wustite (W) that, as previously stated by the same authors [46], seems to ensure complete stability to the slag. In fact, the particle structure appeared unaltered and the phase chemical composition measured by EDS probe pointed out negligible silicon, calcium and iron dissolution (Table 4).

These results highlight the important role of crystalline phases in slag leaching, as previously stated by several authors [6,7,33,34,51–54]. In particular, the stability of wustite was well documented by Belhadj [55], whereas larnite, and its polymorphism bredigite, as well as brownmillerite own hydraulic properties, since they are the same constituents of cement and concrete and behave in the same way when hydrated [36,56,57]. 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, $\text{CaO}\text{-SiO}_2\text{-FeO}_x$ and $\text{CaO}\text{-SiO}_2\text{-Al}_2\text{O}_3$ ternary diagrams. In fact, ternary diagrams provide reliable indications about the crystalline phase/phases in the slag matrix. Referring to the samples showed in Figs. 4 and 5, there is an agreement between their position on ternary diagrams, and XRD and SEM evidences.

Discussion

Barium

The MgO concentration in slag controls the Ba leaching with a quadratic equation (Fig. 6a). If a slag contains enough Mg-complexing oxides (i.e. FeO), barium oxide could dissolve in calcium silicates lattice (i.e. $\beta\text{-Ca}_2\text{SiO}_4$). Unfortunately, this phase has hydraulic properties and dissolves during contact with water. In contrast, in high magnesia slag, MgO could replace barium (and calcium) in calcium silicates, reducing the BaO solubility in such phases (i.e. $\alpha\text{-Ca}_2\text{SiO}_4$) [58,59]. Thus, when Ba is bound in calcium silicates, its release could be enhanced because of the high reactivity of these phases with water. In both cases (with high or low MgO content), Ba could easily be released. In addition, the MgO effect could be divided as a function of its average content in the slag. In fact, as indicated in Fig. 7a, two groups can be identified: the first is characterized by MgO content in the 1–5% wt. range, the latter from 5 to 14% wt. The first group consists most prevalently of slag in groups A and B, whereas the latter is mostly in C group slag.

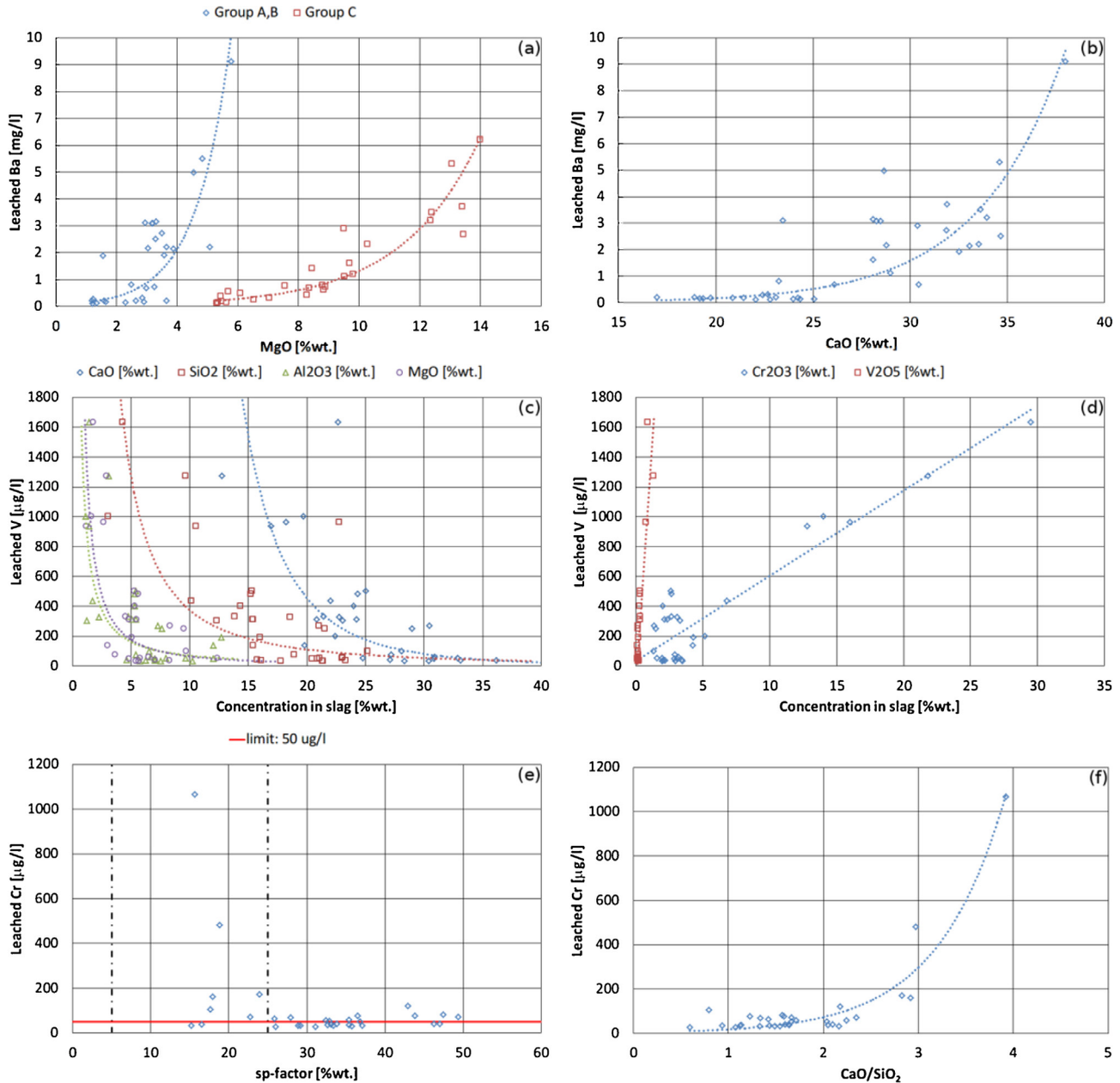


Fig. 6. Effects of slag chemical composition on Ba (a, b), V (c, d) and Cr (e, f) leaching.

Nevertheless, the MgO effect on Ba release is the same for both the groups.

The same conclusions can be reached for the CaO content of the slag. As reported in Fig. 6b, the Ba leaching seems to be favoured by high CaO concentrations. Most likely, high CaO content lead to the formation of high-hydraulic silicates that easily react with water (i.e. Ca_3SiO_5 , Ca_2SiO_4). The hydration process facilitates the migration of species such as Ca, Mg or Ba into water.

The effect of SiO_2 on slag hydraulic behaviour depends on CaO content and for this reason no correlation between SiO_2 and leached Ba was found. Thus, SiO_2 fraction is not a useful index to forecast slag leaching behaviour, and for this reason is not reported. Although several authors [32–34,46] reported the beneficial effect of the addition of silica on retaining behaviour, in this case, the slag analysed was not chemically treated and the silica content is only related to the process route.

Vanadium

V behaviour (Fig. 6c) seemed to be controlled by CaO, SiO_2 , Al_2O_3 and MgO content in the slag. The higher the CaO fraction the lower V concentration there is in the leachate. This result complies with the V-pH relationship and correlates with other research in the same field. Specifically, some yet to be published results have suggested an experimental correlation between Ca and V. Increasing the Ca leaching, V concentration in the leachate solution slightly decreases. V also seemed to have a logarithmic dependence on the aluminium and magnesium oxide weight fractions. A direct correlation between V_2O_5 in the slag and leached V has been found.

Vanadium oxides are usually associated to silicates and calcium-ferrite. Brownmillerite-type ($4\text{CaO}\cdot\text{Al}_2\text{O}_3\cdot\text{Fe}_2\text{O}_3$) phases could also be formed with a low Al_2O_3 amount thanks to the

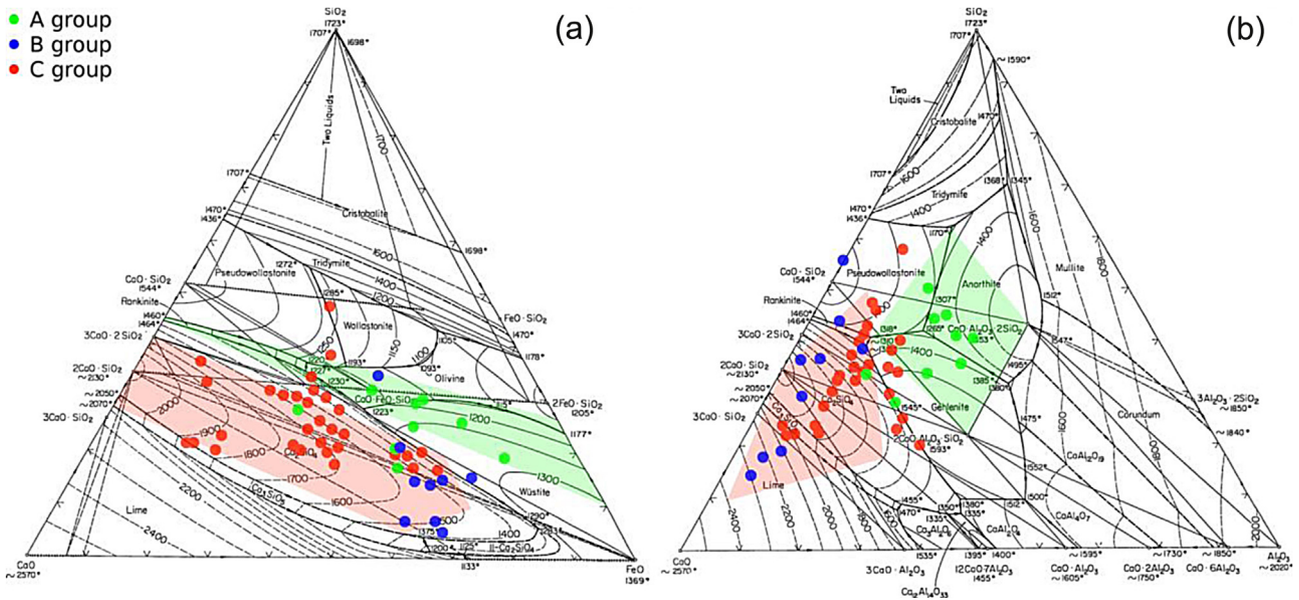


Fig. 7. Identification of safe (green) and risk (red) chemical composition areas on (a) CaO–SiO₂–FeO and (b) CaO–SiO₂–Al₂O₃ ternary diagrams to predict leaching behaviour in standard conditions of EAF carbon steel slag [23]. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

reaction between calcium–ferrite and vanadium and/or chromium oxide. Essentially, SEM-EDS investigations highlighted brownmillerite, constituted with high V and/or Cr oxide in place of Al₂O₃. For example, the brownmillerite identified in Fig. 4 is featured high Cr content (c.a. 9% at.) and non-negligible V content (c.a. 0.5% at.). In other samples, mainly belonging group C, brownmillerite had a composition ranged from 0.5 to 1.0 V and 5–10 Cr (% at.). Since brownmillerite has hydraulic properties and hydrates during the first contact period between slag and water, in poor Al₂O₃ slag, higher V concentration can be eluted. In addition, Al and Mg oxides are the main constituents, together with magnetite, of spinel phases. The analysis carried out on the different slag samples investigated emphasized the vanadium tendency to also be bound in spinel phases. If the slag is featured by low content of MgO and Al₂O₃, non-stable spinels can occur, promoting V leaching. Furthermore, in these conditions, as demonstrated by Kuhn et al. [21,22], Cr can also be easily released.

High MgO content in the slag inhibit V leaching, but enhance Ba leaching. The above mentioned considerations may explain the conflicting behaviour between Ba and V. Effectively, group B slag manifested the release of V but not Ba. Their average chemical composition (reported in Table 3) is characterized by low MgO and Al₂O₃ content, that seem to favour V leaching. On the contrary, the slag characterized by higher Al₂O₃ and MgO content, indicated a higher Ba concentration in the leached solution. Moreover, for the slag featured by a higher CaO fraction, the V concentration in leachate solutions never reached worrying values, with an average concentration of less than 50 µg/l and limited data dispersion. On the other hand, slag featured by a low CaO concentration manifested V leaching, like the other cases. High SiO₂ content in the slag seemed to improve its V-retaining behaviour (Fig. 6c). SiO₂ over the 15% wt. seems to be enough to completely prevent V release. This result confirms the good V-retaining behaviour of acidic slag, as demonstrated in previous work [34,35].

Linear proportional dependence was instead retrieved between V₂O₅, Cr₂O₃ and leached V (Fig. 6d). The higher the Cr₂O₃, the higher the spinel fraction and size in the slag microstructure. High Cr₂O₃ content imposes the presence of higher spinel-forming species (i.e. MgO, Al₂O₃) in order to ensure stable spinel formation [21,22]. If the slag chemical composition, does not allow for the

spinel-forming species involved to be properly combined, unstable spinels are formed, inducing Cr and V leaching. The cooling rate and the cooling procedures also control the spinel formation [12,18], but this type of correlation was not analysed in the present work.

Chromium

Concerning the Cr release behaviour, no indication or clear correlation were found between Cr concentration in the elute and slag chemical composition. Thus, as described by Kuhn et al. [21,22] the main factor ruling Cr immobilization is the formation of spinel structure with composition similar to natural MgCr₂O₄. Even if, nowadays, the only reliable forecasting tool to predict Cr leaching seems to be the *sp-factor* (Eq. (1)) proposed by Kuhn et al. [21,22], its efficacy has not been completely verified for carbon steel slag.

$$sp - factor = 0.2 \times MgO + 1.0 \times Al_2O_3 + n \times FeO_n - 0.5 \times Cr_2O_3 (\%wt.) \quad (1)$$

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.

In fact, as reported in Fig. 6e, several samples that have an *sp-factor* over the 25% wt. (value considered enough to prevent Cr leaching). released Cr, whereas some samples falling within the unsafe zone (between 5 and 25% wt.), did not manifest any Cr leaching. It seems more reliable to focus on the effect of slag basicity (in terms of CaO/SiO₂ ratio) on Cr leaching (Fig. 6f). High slag basicity is associated with a high Cr⁶⁺/Cr³⁺ ratio [60]. Thus, increased slag basicity, decreases the Cr that can be bound in the spinel (in the Cr³⁺ form as Cr₂O₃ [60–62]), increasing the mean chromium content in the slag matrix. In addition, high basicity induces high MgO and CaO activity in the molten slag, resulting in the formation of a Ca-chromite phase instead of a more stable MgCr₂O₄ [63]. Ca-chromite is unstable and its role in Cr leaching is well known [64,65].

The average slag chemical composition could help to predict slag properties, for both the leaching elements and microstructural features, since the relationship between these two aspects is

established. In fact, the dangerous heavy metals involved in the leaching process from EAF carbon steel slag are controlled by the oxide species (i.e. CaO, MgO, . . .) and specific crystalline phases (i.e. larnite, brownmillerite, . . .).

A useful indication about the chemical composition of safe solid slag is indicated in the ternary diagrams shown in Fig. 7. The slag fall in the green areas were featured by stable matrix, mainly formed by gehlenite and/or kirschsteinite and in most of the cases they were compliant with the leaching limits, whereas the slag fall in the red areas were characterized by larnite matrix and subjected to the release of polluting substances. Since the EAF slag can be considered a quinquenary oxide system, the effect of the other oxides species forming the slag (i.e. Al₂O₃ for C–F–S diagram and MgO for C–A–S) can be determined by other pseudo-ternary diagrams for quaternary systems [66–70]. In particular, Al₂O₃ content over 5% by weight creates the formation of gehlenite and anorthite in the CaO–SiO₂–FeO system, closing the CaO–SiO₂ fields. By increasing the Al₂O₃ concentration, however, the olivine field disappears [69,70]. On the contrary, MgO tends to close the gehlenite and anorthite field in the CaO–SiO₂–Al₂O₃ system from 5% by weight. This aspect could explain why slag featured by a high concentration of MgO release high Ba concentration and form gehlenite structure with difficulty [66–68]. For example, the outliers of group A suffered this problem: even if their position on the C–A–S ternary diagram is in the field of gehlenite, their microstructure did not signal gehlenite. On the contrary, the matrix was characterized by high Mg-di-calcium silicate, which induced the high Ba leaching highlighted during the standard leaching test.

Thus a correct balance of chemical composition could allow the formation of stable and safe slag. In particular, MgO should not exceed 5% by weight for high oxidized slag (groups A and B) and the 7% for less oxidized ones (group C), whereas Al₂O₃ should be in the range 7–10% wt. Nonetheless, CaO should be controlled, to avoid reaching an overly high concentration that could stimulate Cr leaching and create dusting and volume instability problems.

Since the slag has to be prepared to obtain steels with the appropriate quality and properties, a thermo-chemical stabilization/inertization process (outside the EAF) should be applied independently on all the recyclable slag in order to avoid dangerous environmental contaminations. In particular, as proposed by Mombelli et al. [51] the addition of suitable external oxides during the deslagging operation could improve the sustainability of the slag, moving the chemical composition in the safe areas of ternary diagrams.

Even if water-slag interaction only interests the slag surface, it is enough to dissolve an important amount of dangerous substances and to drive the leachate pH to caustic values. These results, although qualitative, indicated that liquid-on-solid ratio plays a fundamental role on slag leaching behaviour. The characterization of the slag leaching behaviour through only standard leaching tests could be misleading in order to assess safety or hazard attitude of this by-product. If slags are to be used as unbound stone material, the continuous wetting-drying cycles may lead to dangerous substance accumulation into the environment even if the slag were declared safe after a standard elution test.

These considerations suggest that the characterization of slags should not only undergo the standard elution test but also a modified test in which the effect of liquid-on-solid ratio is investigated.

Conclusions

In this paper different EAF carbon steel slag associated to different steel production were investigated to understand the mechanisms that control the leaching of polluting substances

during leaching tests in standard conditions. The slag chemical composition influences the leaching behaviour of the slag and thus, it can be used to forecast the outcome of leaching test.

On the basis of the obtained results, the following conclusions can be attained:

- slag characterized by high CaO and MgO content (although they indicated caustic leachate pH) had better retaining behaviour against V release. Acidic slag (with high SiO₂ content) was featured by a better trapping attitude against V release;
- high CaO and MgO concentrations lead to high Ba release. A correct balance of the basic oxide species could reduce the slag environmental impact, reducing the overall element leaching;
- high basicity slag, in terms of CaO/SiO₂ ratio, could enhance Cr leaching;
- water-slag interaction interests only the slag surface but was enough to dissolve important amounts of dangerous substances and to drive the leachate pH to caustic values;
- ternary diagrams can be used to predict the leaching behaviour of slag but the effect of other oxides on ternary systems must be taken into account due to their significant effects on phase equilibrium;
- a correct balance of chemical composition could allow the formation of stable and safe slag, eventually providing opportune chemical correction outside the EAF.

In order to satisfactorily reuse EAF slag for civil purposes (aggregates for asphalt, inert gravel, . . .) some practical recommendations can be highlighted: MgO should not exceed 5% by weight for high oxidized slag and the 7% for less oxidized ones, Al₂O₃ should be in the range 7–10% wt. and CaO should not exceed the 30% by weight to fit the safe areas on ternary diagrams.

If the slag chemical composition is out the recommended range, a chemical composition correction by the addition of stabilizers (i.e. SiO₂, Al₂O₃, . . .) would be applied outside the EAF. Slag characterized by stable and non-leachable microstructure can be safely used also in unbound applications, namely unpaved roads, armourstone or gabions.

Appendix A

List of phases.

Phase	Definition	Formula
Akermanite	Calcium-magnesium silicate	Ca ₂ MgSi ₂ O ₇ (2CaO·MgO·2SiO ₂)
Anorthite	Calcium-aluminium silicate	CaAl ₂ Si ₂ O ₈ (CaO·Al ₂ O ₃ ·2SiO ₂)
Bredigite	α di-calcium silicate	Ca ₇ Mg(SiO ₄) ₄ (7CaO·MgO·4SiO ₂)
Brownmillerite	Tetra-calcium aluminate ferrite	Ca ₂ (Al,Fe) ₂ O ₅ (4CaO·Al ₂ O ₃ ·Fe ₂ O ₃)
Gehlenite	Calcium-aluminium silicate	Ca ₂ Al(AlSi) ₂ O ₇ (2CaO·Al ₂ O ₃ ·SiO ₂)
Kirschsteinite	Calcium olivine	CaFeSiO ₄ (CaO·FeO·SiO ₂)
Larnite	β or γ di-calcium silicate	Ca ₂ SiO ₄ (2CaO·SiO ₂)
Mayenite	Calcium aluminate	Ca ₁₂ Al ₁₄ O ₃₃ (12CaO·7Al ₂ O ₃)
Mg–Cr-spinel	Magnesiochromite (spinel)	MgCr ₂ O ₄ /(Mg,Fe)(Al,Cr) ₂ O ₄
Wustite	Iron oxide	(Fe, Mg, Mn)O/MgO·FeO

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