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Hydrothermal Valorization of Steel Slags—Part I: Coupled H2 Production and CO2 Mineral Sequestration

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A new process route for the valorization of BOF steel slags combining H2 production and CO2 mineral sequestration is investigated at 300°C (HT) under hydrothermal conditions. A BOF steel slag stored several weeks outdoor on the production site was used as starting material. To serve as a reference, room temperature (RT) carbonation of the same BOF steel slag has been monitored with in situ Raman spectroscopy and by measuring pH and P\text{CO2} on a time-resolved basis. CO2 uptake under RT and HT are, respectively, 243 and 327 kg CO2/t of fresh steel slag, which add up with the 63 kg of atmospheric CO2 per ton already uptaken by the starting steel slag on the storage site. The CO2 gained by the sample at HT is bounded to the carbonation of brownmillerite. H2 yield decreased by about 30% in comparison to the same experiment performed without added CO2, due to sequestration of ferrous iron in a Mg-rich siderite phase. Ferric iron, initially present in brownmillerite, is partitioned between an Fe-rich clay mineral of saponite type and metastable hematite. Saponite is likely stabilized by the presence of Al, whereas hematite may represent a metastable product of brownmillerite carbonation. Mg-rich wüstite is involved in at least two competing reactions, i.e., oxidation into magnetite and carbonation into siderite. Results of both water-slag and water-CO2-slag experiments after 72 h are consistent with a kinetics enhancement of the former reaction when a CO2 partial pressure imposes a pH between 5 and 6. Three possible valorization routes, (1) RT carbonation prior to hydrothermal oxidation, (2) RT carbonation after hydrothermal treatment, and (3) combined HT carbonation and oxidation are discussed in light of the present results and literature data.

Keywords: hydrogen production, CO2 sequestration, BOF steel slag, hydrothermal oxidation, saponite

INTRODUCTION

In the context of sustainable development and circular economy growth, alternative sources of raw materials represent an ever-expanding market. In the past, a significant part of industrial wastes and byproducts was considered as ultimate waste. Today, waste regulation is evolving toward more recycling. The steel industry is directly concerned by this mind shift. Steel-making processes, which are mastered for long, produce high amounts of by-products. With a production equivalent to 10–15wt.% of the crude steel output (van Oss, 2013) up to 240 millions of tons of steel slags are...
annually produced and only partially valorized. In the last decades, several applications using steel slags have been proposed but only a few of them has been developed at an industrial scale. For example, a small proportion of the slag production is spread on agricultural fields for reclamation of acidic lands and soil stabilization (Chand et al., 2015), taking advantage of a high content in reactive calcium hosted in both lime and portlandite, called “free Ca” in the following. The potassium content also motivated its use as fertilizer (Wang and Cai, 2006).

Some steel slags are incorporated in binder mix for cement and soil stabilization (Chand et al., 2015), taking advantage of its potential on the enhancement of H₂/magnetite partial pressure can potentially meet a double valorization objective.

Based on the literature data cited above, three valorization routes which combine H₂ production and CO₂ sequestration can be envisaged. H₂ can be produced first (Route A, Figure 1), hydrothermal oxidation Reaction (1) will occur at a pH that is imposed by the steel slag chemistry (basically pH > 10). As no other gas is added, H₂ will be produced with high purity and will be compatible with Fuel Cell technologies (Malvoisin et al., 2013) unless the initial slag is significantly carbonated after aging in air.

Then, 2.13 g H₂/kg FeO can be produced after 72 h reaction at 300°C. A later room temperature (RT) aqueous carbonation step can be easily added to sequester CO₂. Aqueous carbonation acts as an accelerated aging and hence improves the residue for its valorization as road engineering product.

A second possible path consists in performing RT aqueous carbonation prior to hydrothermal oxidation (Route B, Figure 1). This path was tested by Malvoisin et al. (2013) and led to a modification of the steel slag reactivity by carbonating the “free Ca.” Hydrothermal oxidation is expected to occur at pH closer to neutral values. A minor effect was reported on the H₂ production yields with the alteration of H₂ purity through the formation of methane for T above 200°C (Malvoisin et al., 2013).

We proposed to investigate here a third route through the application of carbonation and oxidation at once in a single high-temperature step (Route C, Figure 1). Potentially, pH can be dragged to acidic values (pH < 6) through the dissolution of aqueous carbonic acid and drag the pH to acidic values. In addition to its potential on the enhancement of H₂/magnetite production kinetics, CO₂ sequestration will occur coincidentally through carbonation of steel slag minerals. Actually, the carbonation capacities of BOF (or LD) steel slags are high and have been, respectively, estimated, on samples of different origin, to 250 and 142 kg CO₂/t by Huijgen et al. (2005) and Malvoisin et al. (2013), respectively. Therefore, high-temperature oxidation under CO₂ partial pressure can potentially meet a double valorization objective.

In order to better characterize this RedOx reaction, the pure FeO–water system has been investigated by Crouzet et al. (2017) under hydrothermal conditions; FeO oxidation and H₂ production rates were found to be strongly enhanced by replacing pure water by dilute organic acids. For example, Crouzet et al. (2017) showed that FeO is fully oxidized into Fe₃O₄ within less than 10 h at a temperature as low as 150°C using a 0.05 M acetic acid solution (pH of ca. 4.7). In comparison, less than a quarter of the oxidation reaction is achieved after 144 h at 300°C in pure water. However, the kinetics effect of dilute acid obtained on pure FeO cannot be directly transposed to steel slags. Indeed, due to their high Ca content, steel slags will tend to neutralize mild acids and buffer the pH to high values (Crouzet et al., 2017). Therefore, in the perspective of H₂ production from steel slags at an industrial scale, large volumes and/or high concentrations of acid would be required in order to reach significant kinetics enhancement.

In order to meet the objective of enhanced H₂/magnetite production kinetics through acidification of the reacting aqueous medium, we propose here to test the effect of maintaining a CO₂ partial pressure in the course of the hydrothermal treatment of the slag. Indeed, CO₂ partial pressure will buffer the concentration

\[ 3\text{FeO} + \text{H}_2\text{O} = \text{Fe}_3\text{O}_4 + \text{H}_2. \]  

\[ \text{Fe}_3\text{O}_4 \text{ HO} \text{ Fe OH}_2^+ + \text{H}_2. \]  

FIGURE 1 | Three possible routes (A–C) for the hydrothermal valorization of steel slag through H₂ production and CO₂ sequestration. Route c is investigated in the present study.
CO₂ and the formation of carbonates. A kinetics enhancement is then expected (Crouzet et al., 2017).

**MATERIALS AND METHODS**

**Starting Materials**

BOF steel slag was sampled on a French production site. Samples were collected at three different locations on the slag heap where the slag had been dumped, respectively, a few weeks, several months and several years before. The uppermost 10–20 cm were removed and about 30 kg of steel slag were shoveled from the same spot of the heap. In order to increase both reactive surface area and chemical homogeneity, 5 kg of sample were crushed into powder in a jaw crusher and a planetary ball mill. Laser granulometry showed that particles below 50 μm were obtained with sizes centered on 20 μm.

**Ambient Carbonation**

Carbonation of steel slag was performed in a 1 L hastelloy (Ni-Cr alloy) PARR® autoclave equipped with pH measuring device (Montes-Hernández et al., 2009). Ten grams of crushed BOF slag were inserted in 500 mL of deionized water. After dispersion and pH stabilization, carbonation experiment was conducted by injecting 10 bar of CO₂ gas. Pressure was read on a 0–137 bar range manometer. Pressure and pH were recorded during 24 h under 400 rpm stirring. CO₂ pressure drop due to CO₂,aq dissolution was determined separately with the same set-up but without the slag sample. The CO₂ uptake was estimated from the difference between CO₂ pressure drop with and without BOF slags, the steady-state pH in both instances being relatively close, 4 and 3–3.5, respectively. The overall sample carbonation which includes the contribution of carbonates already present in the starting material was determined by thermal gravimetric analysis (TGA). Carbonation extent will be expressed in the form of a CO₂ uptake as the mass of uptake CO₂ (kg) divided by the equivalent mass of the CO₂-free slag (ton) having sequestered that CO₂. It must be noted that the initial slag used here is already partly carbonated. In other words, the slag has already uptaken atmospheric CO₂ into an amount that will be referred to as starting material carbonation (SMC) and which is also calculated on the basis of the mass of CO₂-free starting slag. After a carbonation experiment, TGA data allow to retrieve the overall slag carbonation (OC), i.e., the total amount of CO₂ stored by mass of uncarbonated slag. Note that the difference between OC and SMC yields the effective mass of CO₂ that the slag has already uptaken atmospheric CO₂ into an amount which is also calculated on the basis of the mass of CO₂-free slag. The accuracy on the CO₂ content determination is mostly bounded to the fit of the TGA data and is estimated to better than 5%.

**Solid-Phases Characterization**

Chemical composition (major and trace elements) of the starting materials was determined at CRPG-SARM (Nancy, France) following the procedure described in Carignan et al. (2001). Powdered samples were fused in LiBO₂, dissolved by acid attack and analyzed using flow injection and low pressure on-line liquid chromatography coupled to ICP-MS. Iron (II) was quantified separately. Samples were first dissolved in a sulfuric, hydrofluoric and boric acid mix. Iron (II) amount was then determined by potassium dichromate in phosphoric acid and potentiometric titration. Sample carbonation was characterized by TGA (Metler Toledo TGA/DSC 3+). Between 30 and 50 mg of sample were heated from 25 to 1,100°C at a rate of 10°C/min in N₂ atmosphere. Calcium carbonates from steel slags decompose in the 500–1,000°C range (e.g., Huijgen et al., 2005). Weight loss observed in the 475–800°C range for all studied samples, was therefore assumed to represent CO₂ loss and was used to retrieve carbonation extent. In other words, CO₂ contents derived from TGA here assume CaCO₃ as sole carbonate phase in the sample. Accuracy on the CO₂ content determination is mostly bounded to the fit of the TGA data and is estimated to better than 5%. Total carbon was measured at CRPG-SARM (Nancy) with a CS analyzer (EMIA 320V2 HORIBA) with a precision better than 5%.

Mineral phases were identified by X-ray powder diffraction (XRPD) with a D8 diffractometer (Bruker, CuKα radiation) operated with a 2θ step size of 0.026°. Quantitative analysis of crystalline phases was conducted on the HT-carbonated sample by Rietveld refinement using the BGMN software (Taut et al., 1998). Attempts on the two other samples (starting and RT-carbonated slag) were not conclusive due to the likely presence of amorphous to poorly crystalline phases.

**Hydrothermal H₂ Production**

Hydrothermal slag oxidation was conducted in a third 500 mL Hastelloy® autoclave (Crouzet et al., 2017) under 500 rpm stirring. Four grams of steel slag were inserted in 200 mL of deionized water. About 10 bar of CO₂ were injected at ambient temperature. After 2 h, Pₐ total stabilized after it dropped by about 6 bar, argon was then injected to reach 100 bar of total pressure (Pₐ total). Pₐ total was then lowered by 50 bar in order to reach 190 bar at 300°C. Gas phase was sampled at regular intervals to monitor hydrogen and other gases (CO₂, N₂, O₂, CO, and CH₄). Gas was sampled directly in the hot autoclave, water was condensed and the dried gas readily analyzed with a Clarus 500 gas chromatography (GC). The GC is equipped with a thermal conductivity detector and a Restek ShinCarbon® column, polymer filled and 2 m long. Argon was used as gas carrier. The temperature of the detector, the injection system and the oven were, respectively, set to 250, 100, and 80°C. Each gas sample was analyzed at least three times consecutively. H₂ concentration in the sampled gas is converted into absolute quantity (e.g., mass) in the reactor using the ideal gas law. Mass of H₂ is normalized to the equivalent mass of FeO included in the solid starting material. Initial steel slag FeO contents were deduced from chemical analysis (Table 1).
Solid samples were also observed by field-emission scanning electron microscopy (FE-SEM) on a ZEISS Ultra 55 equipped with an energy dispersive X-ray spectroscopy (EDS) detector. Two preparations were employed prior to carbon coating. For chemical analysis, the sample was embedded in epoxy and polished. Crystal morphologies were observed on powder sample dispersed on a double-sided carbon tape.

**RESULTS**

**Starting Steel-Slag Characterization**

**Chemical Composition**

Composition of the three starting steel slags was measured by ICP-AES and -MS analysis. Average results from three replicas are presented in Table 1. Accounting for nearly half of the sample mass, calcium oxide is the main constituent of the steel slag samples followed by iron oxide and silica. Average chemical compositions are in good agreement with literature values (Proctor et al., 2000; Malvoisin et al., 2013; Piatak et al., 2015). Only minor variations in chemical composition were observed between the three samples, this chemical homogeneity among samples reflects the Basic Oxygen Furnace process mastering. Considering that CO₂ sequestration occurs through calcium carbonate formation, an overall carbonation potential of 387 kg CO₂ can be deduced from the bulk CaO content. SMC was determined to be 63 kg CO₂/t steel slag on the freshest sample (weeks, Table 1) by TGA (Figure 2). This SMC figure corresponds to 5.9 wt.% CO₂ in the slag (Table 1).

Total CO₂, LOI, and iron oxidation are the chemical parameters that significantly vary with storage time (Table 1). The longer the steel slag was stored, the higher the total CO₂ content. Iron oxidation due to weathering is evidenced by a decrease of the Fe²⁺/ΣFe ratio as storage time increases. This ratio drifted from 45% after weeks and months to 33% after years of outdoor storage.

**Starting Steel Slag Mineralogy**

Mineralogical content of the BOF slag sample aged for several weeks on the storage site (Figure 3) was determined combining bulk composition (Table 1), phase identification by XRPD, FE-SEM imaging, and EDS analysis (Table 2; Figure 4). Three calcium carbonates were identified by XRPD: calcite, aragonite, and vaterite (Table 2). Based on TGA measurement which yielded 5.9 wt.% CO₂, these calcium carbonates represent around 15 wt.% of the initial steel slag (Figure 2). Larnite, Ca₂SiO₄ (C₂S), and brownmillerite are the main mineral phases in the starting slag which account together for more than 50% of the slag mass. Brownmillerite and magnetite host the ferric iron contained in the starting steel slag. Ferrous iron is hosted in magnesian wüstite (called wüstite for simplification hereafter).

**RT Carbonation**

**Time-Resolved Carbonation: pH, P_CO₂ Drop, and In Situ Raman Monitoring**

Carbonation was conducted at RT on the freshest slag immersed in water. CO₂ pressure drop and pH were first monitored over
Table 2: Mineralogical characterization of both starting material and experimental products.

<table>
<thead>
<tr>
<th>Mineral phases (XRPD + SEM-EDS)</th>
<th>Initial steel slag (weeks)</th>
<th>RT carbonated (25°C)</th>
<th>HT carbonated (300°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Main mineral phases</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium (CaCO₃)</td>
<td></td>
<td>Calcite</td>
<td>Calcrete [66.0(7)],</td>
</tr>
<tr>
<td>Larnite (Ca₂SiO₄)</td>
<td></td>
<td>brownmillerite</td>
<td>siderite [10.2(3)],</td>
</tr>
<tr>
<td>Brownmillerite (Ca₂Fe₁.₅Al₀.₅O₅)</td>
<td></td>
<td>wüstite</td>
<td>saponite [11.7(8)],</td>
</tr>
<tr>
<td>Wüstite (Fe₀.₅₂Mg₀.₄₈O)</td>
<td></td>
<td></td>
<td>magnetite [5.7(3)]</td>
</tr>
<tr>
<td>Portlandite (Ca(OH)₂)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aragonite (CaCO₃)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minor phases (&lt;5 wt.%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vaterite (Ca₃(CO₃)₂)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OH-apatite (Ca₅(PO₄)₃OH)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OH-apatite magnetite</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>OH-apatite hematite</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magnetite (Fe₃O₄)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CO₂ uptake (kg/ton of fresh slag)</td>
<td>63 (SMC)</td>
<td>306 (OC)</td>
<td>390 (OC)</td>
</tr>
</tbody>
</table>

Mineral phases are divided into two classes depending on their estimated abundance using semiquantitative XRPD. Composition in parentheses is derived from EDS data on polished samples. For the hydrothermally carbonated sample (HT carbonated), crystallinity of the phases allowed reliable Rietveld refinement, mineral weight proportions are given in brackets with the SD. Note that OH-apatite has only been unequivocally identified in the HT-carbonated sample, another Ca-phosphate might be present in the two other samples but has been assumed to be OH-apatite. CO₂ uptake is determined using TGA, SMC, and OC stand for starting material carbonation and overall carbonation, EC (experimental carbonation, i.e., CO₂ uptake in the course of the carbonation experiment) is calculated by subtracting SMC (initial steel slag) to OC for both RT and HT-carbonated samples.

*See text for discussion about saponite composition.

In parallel to CO₂-pressure and pH monitoring, time-resolved phase changes in the aqueous suspension were monitored by in situ Raman spectroscopy (Figures 5C,D). Before CO₂ injection, portlandite and calcite were the two main components.
**FIGURE 4** | Mineralogical content of the starting steel slag: (A) X-ray powder diffraction (XRPD) data: portlandite (P), calcite (C), larnite (L), brownmillerite (B), and wüstite (W); (B) field-emission scanning electron microscopy (FE-SEM) image in back-scattered electron mode: (1) (Fe_{0.52}Mg_{0.48})O wüstite, (2) (Fe_{0.45}Mg_{0.55})O wüstite, and (3) intergrowth: 85 mol.% Ca_2SiO_4 + 15 mol.% of Ca-phosphate assumed to be Ca_5(PO_4)_3OH.

**FIGURE 5** | Time-resolved monitoring of steel slag carbonation at ambient temperature. (A) CO₂ pressure drop in water (squares) and with the steel slag (circles). Dotted lines correspond to a fit to the equation: \( P_{\text{CO}_2, \text{diss}} = P_{\text{CO}_2, \text{initial}} - A_{\text{CO}_2} \times e^{-kt} \). (B) Time-resolved pH data fitted with three parameters: \( \text{pH}(t) = A_{\text{pH}} \times e^{-kt} + B_{\text{pH}} \). (C-D) In situ Raman spectra presented in two wavenumber ranges.
identified in the suspension. After CO₂ was injected, portlandite consumption started and lasted until its total disappearance after 20 min of reaction. Calcite signal started to grow after 10 min of reaction. It kept growing even after portlandite consumption until 720 min of reaction and the observation of signal instabilities. In addition to calcite growth, a minor peak belonging to aragonite was transiently detected at 210 cm⁻¹, which disappeared after 420 min of reaction.

Figure 5D shows spectra in the 1,200–600 cm⁻¹ wavenumber range for five different run durations. Calcite is easily identified with two characteristic bands, one at 1,095 cm⁻¹ and another at 715 cm⁻¹. As reaction proceeds, a narrowing of the Raman band in the 710–680 cm⁻¹ range is observed after more than 30 min of reaction. This signal is attributed to the respective contributions of portlandite and a calcium silicate, possibly larnite. At t₀ another small contribution is also detected at 960 cm⁻¹ which quickly disappeared after 20 min of carbonation. This contribution may be related to the partial hydration product of calcium silicate such as afwillite or hibschite. Signal instabilities were observed after 720 min of reaction. As the sapphire peak from the window remains constant, these instabilities are not related to the Raman light source. After opening the autoclave, most of the solid sample was found to have formed a hardened paste, stuck at the bottom of the autoclave, i.e., away from the volume sampled by the Raman probe.

Solid Characterization
X-ray powder diffraction patterns confirmed the full consumption of portlandite and calcium silicates (larnite) as well as the crystallization of calcite (Table 2). Starting aragonite and vaterite all converted into calcite. TGA data indicate a single weight loss event (ca. 23 wt.%) consistent with calcite as the sole Ca-carbonate in the system (Figure 2) and which is characterized by a DTA peak centered around 765°C. An OC of 306 kg CO₂/t is deduced which corresponds to an experimental carbonation (EC) of 243 kg CO₂/t steel slag. It is calculated that about 79% of the CaO mass in the slag is now combined to CO₂ in the form of carbonates. In the starting steel slag (Figure 3), CaCO₃, larnite, and portlandite hosted about 70% of the CaO mass contained in the sample. Therefore, the derived OC is broadly consistent with the carbonation of portlandite and larnite although partial carbonation of brownmillerite cannot be excluded. No major mineralogical changes were observed among iron-bearing compounds. Iron was mostly recovered as oxides in the form of brownmillerite, magnetite, and wüstite (Table 2).

Steel Slag Hydrothermal Oxidation under CO₂ Pressure

Hydrogen Production
Steel slag hydrothermal oxidation was performed at 300°C according to two experiments, i.e., with and without CO₂ (Figure 6). Hydrogen was produced in both cases following a similar kinetic model which is satisfactorily fitted to a simple squared root function of time:

$$H_2(t) = A \cdot t^{1/2}.$$  \hspace{1cm} (2)

In the absence of added CO₂, hydrogen production rate was found to be roughly constant during the 30 first hours before
it progressively slowed down. A total of 2.13 g H$_2$/kg FeO were produced after 72 h and $A_1$ was estimated to 0.276 g H$_2$/h$^{1/2}$/kg FeO. In the presence of CO$_2$ (initial $P_{CO_2} = 2$ bar), a total of 1.55 g H$_2$/kg FeO was produced after 72 h; $A_2$ in Eq. 2 is estimated to 0.185 g H$_2$/h$^{1/2}$/kg FeO. CO$_2$ gas concentration was found roughly constant over the whole reaction. It corresponded to an average CO$_2$ partial pressure of 16 bar at 300°C as calculated assuming CO$_2$ as an ideal gas. No CH$_4$ was detected in the analyzed gas with a CH$_4$ detection limit estimated at 100 ppm.

**Characterization of the HT Carbonation Mineral Products**

As for RT carbonation, calcite is the dominant mineral phase in the sample carbonated at 300°C (Table 2). XRPD (Figure 7A) and SEM characterization (Figure 7B) revealed another carbonate with a composition (Fe$_{0.56}$Mg$_{0.36}$Mn$_{0.06}$Ca$_{0.02}$)CO$_3$ as inferred from EDS data. The diffraction signal of this carbonate could be refined using a siderite structure assuming however lattice parameters that significantly exceed those expected along the siderite-magnesite solid solution. XRPD pattern of HT-carbonated products also revealed the crystallization of saponite, a trioctahedral clay mineral which belongs to the smectite group. The retrieved lattice parameter, $b$, of around 9.22 Å, suggests that the newly formed saponite has octahedral sites dominated by iron (Brigatti, 1983), Fe$^{2+}$ (Baldermann et al., 2014), and/or Fe$^{3+}$ (e.g., Treiman et al., 2014). Iron mass balance from Rietveld data along with the constraint that Fe$^{2+}$/Fe$_{tot}$ in the slag must decrease in order to account for measured H$_2$ production, suggesting that Fe in saponite is dominantly ferric.

Thermal gravimetric analysis data indicate a two-stage mass loss (Figure 2) with two DTA peaks located at 530 and 789°C. From 120 to 1,000°C, the sample has lost about 30% of its initial mass, whereas Rietveld refinement predicts a (H$_2$O + CO$_2$) content in the HT-carbonated slag of 35.5 wt.%. The main mass loss event (24.2 wt.%) corresponds to the 789°C DTA peak and is very likely correlated with calcite decomposition since

---

**FIGURE 7** | X-ray powder diffraction (XRPD) pattern and field-emission scanning electron microscopy (FE-SEM) imaging of the HT-carbonated steel slag; (A) XRPD data: calcite as (C,*), siderite (S,*), clay minerals (Cl), magnetite (M), and hematite (H); (B) BSE mode imaging on polished sample: EDS chemical analysis (1) of Mg-rich siderite (Fe$_{0.56}$Mg$_{0.36}$Mn$_{0.06}$Ca$_{0.02}$)CO$_3$; secondary electron mode imaging of (C) micrometric euhedral calcium carbonates and (D) iron oxide nanoparticles in the 10-30 nm size range.
calcite is the dominant mineral according to XRPD. The lower temperature mass loss (ca. 4 wt.%) may be correlated with either Fe-rich saponite dehydroxylation (Brigatti, 1983; Földvári, 2011) or Mg-siderite decomposition since 530°C is comprised between magnesite and siderite TGA decomposition temperatures (Földvári, 2011). Based on the amount of Fe-saponite (11 wt.%) obtained from Rietveld refinement, the contribution of saponite dehydroxylation to the sample weight loss should not exceed half a weight percent and therefore the lower temperature TGA peak is more likely related to Mg-siderite decomposition. A 3.5–4% mass loss would correspond to the decomposition of 8–9 wt.% siderite which compares relatively well with the 10 wt.% inferred from Rietveld refinement. EC is estimated to 327 kg CO₂/t from TGA. Moreover, no more wüstite nor brownmillerite was found in the XRPD pattern. Iron is now partitioned between magnetite, hematite, saponite and siderite. Carbonates were observed in the form of rhombohedral micrometric particles (Figure 7C).

Further FE-SEM characterization was performed on iron oxides extracted magnetically with a permanent magnet in a water-filled ultrasonic bath. This magnetic separation allowed to emphasize a multitude of iron oxide nanoparticles (Figure 7D).

**DISCUSSION**

**RT Carbonation: Time-resolved Monitoring**

As shown by previous studies on steel slags and other Ca-rich materials at RT (e.g., Huijgen et al., 2005; Montes-Hernandez et al., 2009; Fritz et al., 2013), aqueous carbonation leads to a pH shift from basic to acidic conditions. We find for the steel slag studied here a pH decrease from 10.6 to 3.7 in less than 15 min. From 15 to 40 min and until the end of experiment, pH stabilizes from 3.7 to 3.3. In the Ca(OH)₂–water subsystem, portlandite consumption is enslaved to the supply of protons from CO₂ dissolution; variation in the ionic activity product leads to the very rapid consumption of portlandite and a pH drop by 5–7 units (Fritz et al., 2013). In our experiments with steel slag, a first similar stage was observed but limited to the first minute. This stage may be shortened due to the steel slag chemistry and the availability of ionic charges, supplied by the partial dissolution of other solids either amorphous or crystalline.

*In situ* Raman spectroscopy was used to follow up the mineralogy of the suspension. Before room-temperature (RT) carbonation, portlandite and calcite were the two main mineral species. The first 20 min showed the crystallization of calcite at the expense of portlandite which fully disappeared according to the overall reaction:

\[
\text{Ca(OH)₂(s) + CO₂(g) → CaCO₃(s) + H₂O(l).} \tag{3}
\]

As CO₂ is in excess, this reaction only stops when portlandite is consumed. *In situ* pH measurement shows that the main pH drop occurs during these first 20 min. Despite the persistence of portlandite in the first 20 min of reaction, pH cannot be maintained to high values; it follows that portlandite dissolution is slower than CaCO₃ formation. It must also be noted that metastable carbonates (aragonite and vaterite) which likely formed by interaction between the slag and atmospheric CO₂ converted into calcite in a CO₂-rich aqueous medium at RT (Ogino et al., 1987). Aragonite was even shown to grow metastably in the first hours of reaction before it finally disappeared from Raman spectra after 7 h and was no longer found in the XRPD pattern of the recovered solid sample.

\[ \text{P}_{\text{CO₂}} \] has been used as a third proxy of the carbonation kinetics. We showed that 83% of the total \( \text{P}_{\text{CO₂}} \) drop occurred in the first 85 min and almost no further variation was observed after 407 min. CO₂ uptake from the gas phase lasted however longer than the single portlandite-carbonation event which is estimated to less than 20 min according to our *in situ* Raman spectroscopy data. Following the results by Huijgen et al. (2005), it can be argued that calcium silicates and Ca-bearing oxides carbonation may still proceed after portlandite consumption. This assumption is corroborated by further calcite growth after portlandite disappearance as observed by Raman spectroscopy. At the end of RT carbonation experiment, steel slag is recovered in the form of a hardened paste stuck at the bottom of the autoclave. This hardening process may be enhanced by the cementitious properties of calcium carbonates (Johnson et al., 2003) and of the products of dicalcium silicate (C₂S) hydration.

Based on the CaO content of 49.3 wt.% in the fresh slag (i.e., CO₂-free reference) and on the relationship between CO₂ content and carbonation degree (Relation 6 in Huijgen et al., 2005) for a carbonation degree of 100%, the maximum CO₂ sequestration (maximum OC) by our steel slag sample is estimated to 387 kg CO₂/t of fresh slag. After carbonation at RT, an OC of 306 kg CO₂/t of fresh slag is attained corresponding to a carbonation degree of around 80%, i.e., 80% of the Ca contained in the starting material has reacted to form calcium carbonate. The XRPD pattern of the carbonated slag shows that the unreacted Ca is bounded to brownmillerite which appears to be the least reactive Ca-bearing mineral in the starting material.

**High-Temperature Steel Slag Carbonation**

**Influence of CO₂ Partial Pressure on Hydrogen Production**

In order to investigate the potential of the CO₂ injection at high temperature in lowering the pH and enhancing oxidation rate (H₂ production), hydrothermal steel slag oxidation was performed in deionized water at 300°C to 150 bar under argon atmosphere with and without added CO₂ partial pressure. Hydrogen production was observed in both cases, following similar kinetic models. So apparently, the addition of CO₂ does not fundamentally modify the steel slag oxidation process. In absence of a plateau in H₂ production, a square root function of time was found to yield a better fit to the H₂ production data than a first-order kinetic model (Crouzet et al., 2017). These data indicate a H₂ yield by 30% lower when CO₂ gas is added. We also show that the addition of CO₂ gas leads to the disappearance of wüstite which is otherwise observed as residual phase.
There is a clear discrepancy between recovered H₂ and wüstite consumption in the HT carbonation experiments. This apparent H₂ deficit can be explained by the formation of an (Fe,Mg)-carbonate, \((\text{Fe}_{1-x}\text{Mg}_x\text{Mn}_{0.3}\text{Ca}_{0.1})\text{CO}_3\), of siderite structure (Rietveld refinement) which actually incorporated ferrous iron and prevented it from further oxidation. The Fe/Mg ratio of this carbonate as well as the presence of manganese traces suggest that it directly formed from wüstite with little fractionation of the divalent cations (Fe, Mg, Mn). Another remarkable feature is the presence of hematite. Using the concentration of aqueous H₂ at the magnetite–hematite equilibrium and the Henry constant of H₂ as provided by SUPCRT92 (Johnson et al., 1992), it is calculated that, at 300°C and 190 bar, hematite should no longer be stable over 1% reaction progress (Appendix). Hematite might therefore be an intermediate compound of brownmillerite breakdown which persists as a metastable phase indicating a relatively low reduction kinetics into magnetite.

Total H₂ produced in the carbonation experiment was found to only represent 37 mol.% of maximum H₂ that can be produced based on the initial Fe⁺⁺ content of the starting slag. On the other hand, about 50 wt.% of the initial FeO content is sequestered in iron carbonates. Furthermore, the reduction of hematite into magnetite might consume H₂ and thereby decrease the overall H₂ yield.

Assuming that the pH in the HT carbonation experiments is controlled by the calcite–CO₂ equilibrium, an estimated pH of ca. 5.6 is calculated for the investigated experimental conditions. Crouzet et al. (2017) noted a significant kinetic improvement in FeO oxidation in the presence of acetic acid solution at a pH of ca. 4.7. Unfortunately, in the present HT carbonation experiments, slag wüstite is involved in two competing reactions, oxidation on one hand and carbonation on the other. It is therefore not possible to precisely assess the effect of shifting the pH below 7 on wüstite oxidation kinetics alone. However, in the 72-h carbonation/oxidation experiment, 1.55 g H₂/kg FeO has been produced. If now, we consider that about half of the FeO has been entrapped into carbonates (additional Fe⁺⁺ might also have been incorporated into saponite), it follows that more than 3 g H₂/kg of available FeO has been effectively produced in the time span, i.e., 40% more than in the hydrothermal experiment where no CO₂ was added. It remains that HT carbonation lowered the H₂ yield in comparison to wüstite oxidation in water without added CO₂. Moreover, injection of CO₂ gas for the sake of kinetic improvements led to the alteration of hydrogen purity. As far as H₂ production is concerned slag oxidation under P_{CO₂} has obviously no added value.

In parallel to H₂ production, hydrothermal oxidation/carbonation of steel slag led to the formation of magnetite. FE-SEM investigations were conducted on magnetically separated samples. Magnetite was observed as nanoparticle agglomerates. Crouzet et al. (2017) already reported the formation of magnetite nanoparticles as the result of the hydrothermal oxidation of FeO in the presence of acetic acid solution. Magnetite nanoparticles and its economical relevance are discussed in the companion article (Crouzet et al., in press). It would however be interesting to study whether HT carbonation could influence the properties of the produced nanomagnetite (size, morphology, and composition).

Other H₂ production routes using CO₂ still need to be explored. For example, recently, Michiels et al. (2015) showed that the in situ formation of aqueous CO\(_2\)⁻ has a catalytic effect on the mild hydrothermal oxidation of iron metal powder. The stability of aqueous CO\(_2\)⁻ required these experiments to be conducted at pH = 11 with potassium hydroxide. This catalytic effect was found to be dependent of the CO₂ partial pressure. For an initial CO₂ overpressure of 15 bar in the reactor at 25°C, the authors reported that siderite formation limits H₂/magnetite yield in the 140–260°C range. If this result on Fe metal can be transposed to FeO, slag oxidation under P_{CO₂} at pH > 12 may be worth exploring.

**Improvement of Mineral CO₂ Sequestration at HT**

Although HT carbonation occurred to be detrimental to the H₂ yield, it led to higher carbonation degree compared to RT carbonation. Indeed, mineral sequestration (OC) is, respectively, estimated to 306 and 390 kg CO₂/t of fresh steel slag at 30 and 300°C by TGA (Table 2). Note that the carbonation degree obtained at HT exceeds 100% as defined on the basis of the Ca-content of the steel slag used in this study. This CO₂ sequestration gain at 300°C is explained by (1) the partial carbonation of magnesio-wüstite and (2) the carbonation of the calcium contained in brownmillerite at high temperature. Indeed, no more brownmillerite was detected in the sample recovered from HT carbonation, whereas brownmillerite is still present in the RT-carbonated one. Brownmillerite carbonation liberates Al and Fe\(^{3+}\) which are likely responsible for the formation of saponite and hematite, respectively, as secondary products.

**Toward a New Valorization Route?**

In addition to valorization routes that would involve room-temperature CO₂ sequestration either before or after H₂ hydrothermal production, we investigated a third route through the application of carbonation and oxidation, at once, in a single high temperature step (Figure 1C). Obviously, the main drawback of this last route is that the purity of hydrogen will be strongly altered by dilution in a CO₂-rich gas. Furthermore, contrary to our expectations and even if acidic conditions were reached, H₂ yield decreased by 30% when a CO₂ partial pressure (around 15 bar) was maintained at HT. The only tangible advantage is the enhanced CO₂ sequestration degree of 390 kg CO₂/t of fresh steel slag obtained at 300°C instead of 306 kg CO₂/t at RT. If this valorization route is apparently not promising, one should bear in mind that the economic value of the proposed route lies on the nanomagnetite (by)product (Crouzet et al., in press). Despite a magnetite yield which will be lowered as for H₂ to which Fe₃O₄ formation is coupled, the effect of combining HT oxidation and carbonation on the properties of the produced magnetite (size, morphology, surface, etc...) remains to be evaluated.

**AUTHOR CONTRIBUTIONS**

CC performed all experiments and wrote most of the manuscript. FB supervised the overall work and significantly contributed to the writing of the manuscript. GM-H designed and supervised the carbonation experiments. NR brought is knowledge of RedOx...
reaction. NF carried out XRPD and contributed to the interpretation of the diffraction data. J-HF helped contributed to the definition of industrial applications. BG provided the knowledge of magnetite applications and market.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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APPENDIX

Stability limit of hematite is calculated using the two following equilibria and their respective pK calculated with SUPCRT92 (Johnson et al., 1992):

\[
3 \text{Fe}_2\text{O}_3 + \text{H}_{2,\text{aq}} = 2 \text{Fe}_3\text{O}_4 + \text{H}_2\text{O} \quad (pK_1 = -4.70)
\]

\[
\text{H}_{2,\text{aq}} = \text{H}_{2,\text{g}} \quad (pK_{HC} = -2.19).
\]

Assuming that both solution and gas volumes in the autoclave remain constant between ambient and 300°C–190 bar, i.e., 200 and 300 mL, respectively, and neglecting the initial presence of O₂, it is calculated that ca. 20 μmol of H₂ are produced to reach hematite/magnetite equilibrium. Four grams of slag containing 10 wt.% FeO can produce 1.8 × 10⁻³ mol H₂ at most. Then, the minimum of 20 μmol of H₂ which is required to drive hematite reduction will be produced after a reaction progress of ca. 1–1.5%.