



# Insights Into Nickel Slag Carbonation in a Stirred Bead Mill

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This work is part of the ongoing development of the attrition-leaching carbonation process, a single-step aqueous carbonation technology that integrates a stirred bead mill. The principle of the attrition-leaching carbonation process is to continuously refresh the surfaces of reactive particles so that leaching can proceed unimpeded, yielding enhanced carbonation kinetics and yield. Invariably, attrition-leaching carbonation experiments carried out under controlled temperature and CO<sub>2</sub> partial pressure conditions with different silicate-rich carbonation feedstocks - natural ores and nickel slags - show a carbonation yield that tends towards a plateau 20-50% below the stoichiometric yield. This repeatable behaviour raises the question as to whether the carbonation limitation is due to specific equilibrium side reactions of the carbonation feedstock or to a kinetic limitation of the attrition-leaching carbonation process. To try to provide some answers to this puzzling question, this reflexive paper implements a "thermo-kinetic" modelling methodology based upon geochemical equilibrium simulations and particle reaction models. The results obtained indicate that the observed slowing down of the carbonation process can be explained either by the formation of Mg- and/or Fe-rich silicates that precipitate at the expense of carbonates, or by a decrease in the efficiency of the attrition process over time. Indeed, either one of these mechanisms could explain the observed behaviour of the attrition-leaching carbonation process. However, the cross comparison of different data sources pleads in favour of the attrition-leaching carbonation performance being limited by the attrition process. Pending further confirmation, this tentative conclusion suggests that further development of the attrition-leaching carbonation process requires new knowledge about its inner workings, and that there may be ways to optimize the performance of this process beyond that based on standard stirred bead mill operating rules.

Keywords: aqueous mineral carbonation, slag valorization, attrition-leaching carbonation process, thermo-kinetic modelling, shrinking particle model, shrinking core model

# INTRODUCTION

For several years now, aqueous mineral carbonation has stood out not only as a viable solution for  $CO_2$  sequestration in the form of stable carbonates, but also as a way of treating or recovering littleexploited minerals or mining industry wastes (Huijgen et al., 2005; Sipilä et al., 2008; Bodor et al., 2013; Sanna and Maroto-Valer, 2017; Veetil and Hitch, 2020). This process indeed allows the passivation or dilution of hazardous substances such as or asbestos or heavy metals (Gadikota et al., 2014), the feedstock beneficiation as aggregates or fillers (for road construction, concrete making, soil

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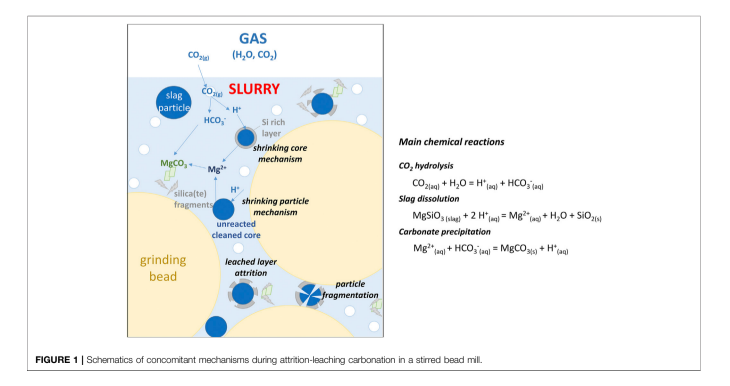
improvement, etc.) or, even better, for the synthesis of building materials with improved mechanical properties and reduced footprint (National carbon Academies of Sciences, Engineering, and Medicine, 2019). The Carboscories/Carboval projects (Bourgeois et al., 2020) aim to contribute to the latter by valorising slags from New Caledonias nickel industry as mineral additions and hydraulic binders. While their chemical composition is rather uniform from one industrial source to another, nickel slags exhibit different mineralogy depending on their cooling conditions. For instance, the air-dried carbonation feedstock considered in the present work is mainly crystallized magnesium silicates with iron and aluminium impurities, while other slags obtained from water quenching are mainly composed of vitreous phases. Not only does the mineralogy control the aqueous solubility of the slag, but it also affects the nature of the solid products that may form during carbonation at the expense of the desirable ones, namely carbonates and amorphous silica. Moreover, solid products (through dissolution/precipitation mechanisms) precipitate on the surface of the slag particles and form surface layers of variable thickness and permeability that limit the accessibility of the protons to the unreacted core and yield low conversion rates. The attrition-leaching carbonation process (Julcour et al., 2015; Bourgeois et al., 2020) has made it possible to remove all or part of these diffusion barriers, enabling the conversion of more than 50% of carbonatable species in about 10 h. This process uses the direct aqueous carbonation route, whose chemistry relies on two main features: 1) gaseous CO<sub>2</sub> is the only reactant; 2) CO<sub>2</sub> hydrolysis provides the proton source for the dissolution of the solid phase in water (see Figure 1). Attrition-leaching carbonation is operated inside a stirred bead mill, a proven industrial process for producing ultrafine particles (Jankovic, 2000). Optimization of the attrition-leaching

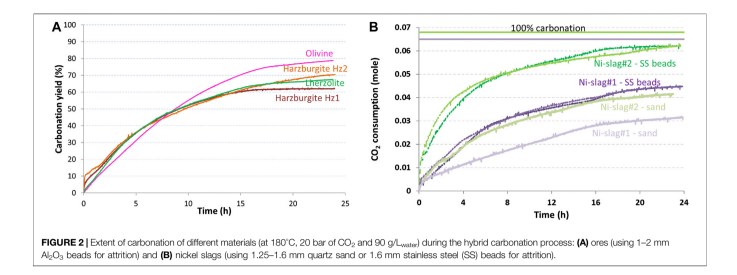
carbonation process is related to matching surface layer formation and attrition rates. To this end, a better understanding of the process features is required. This is actually a complex system with several concomitant mechanisms (see **Figure 1**): attrition, but probably also particle breakage, chemical attack of the reactive feedstock phases, gas dissolution and precipitation of product phases.

## PROBLEM STATEMENT

The authors have envisioned the attrition-leaching carbonation as a process that can be tuned to reach the thermodynamic carbonation limits of the carbonation feedstock, notwithstanding the kinetics involved with reaching this limit. As shown in **Figure 2A** for natural ores (which are all ferromagnesian silicates with various degrees of serpentinisation) and in **Figure 2B** for nickel slags, they have shown that attritionleaching carbonation experiments yield largely material insensitive carbonation kinetics under nearly identical grinding conditions (same agitator and agitation speed, 180°C, 20 bar of  $CO_2$ ). This observation confirms the robustness of the attritionleaching carbonation process, keeping in mind that carbonation yield would not exceed 10% with these materials should the attrition component not be coupled with the reaction.

However, invariably, the carbonation curves tend towards a plateau that is 20–50% below the stoichiometric carbonation yield. These observations lead to the main question that lies behind this paper: is this early slowing down in carbonation extent due to a thermodynamic limit linked with the carbonation feedstocks, or is it a intrinsic feature of the attrition-leaching carbonation process, or both?





Moreover, **Figure 2B** shows that grinding medium selection, all operating conditions being equal, yields different carbonation plateaus. Stirred mill operating conditions, and grinding medium selection in particular, are thus clearly part of the equation. Bringing elements of answer to this consistent behaviour of attrition-leaching carbonation in a stirred bead mill is the objective of this reflexive paper.

The paper uses a "thermo-kinetic" modelling methodology, under the assumption that both gas dissolution and precipitation of product phases are rapid phenomena compared to leaching and attrition processes. The thermodynamic component of the methodology uses geochemical equilibrium modelling to predict the phases that can form under given temperature and pressure conditions (Cassayre et al., 2016). The kinetic component uses traditional shrinking core and/or shrinking particle models for reacting particles. Predictions are assessed against experimental data obtained in a 300 ml laboratory stirred mill with a fully crystallized pyroxene-rich nickel slag whose leaching kinetics can be compared to published data (Oelkers and Schott, 2001).

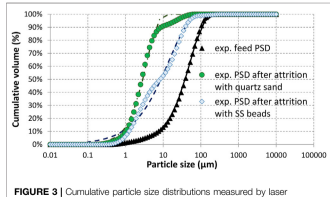
# MATERIALS AND METHODS

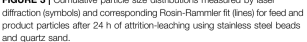
### **Processed Material**

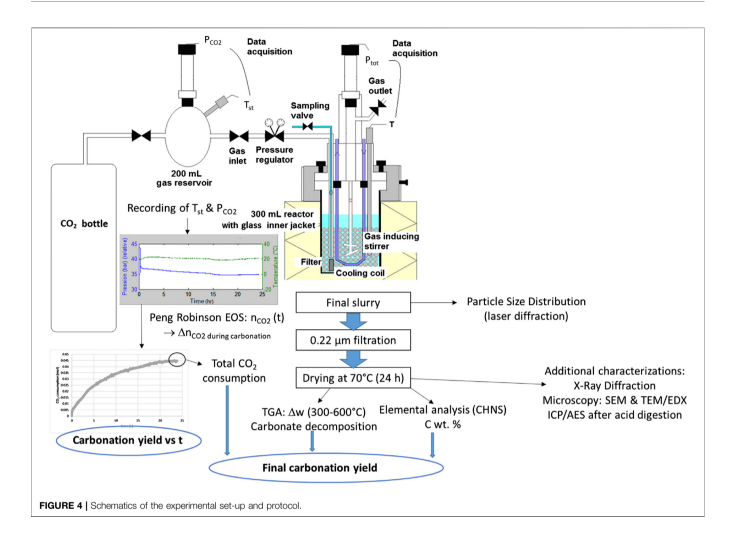
The carbonation feedstock is a magnesium silicate slag from a nickel pyrometallurgical plant that operates in New-Caledonia (Koniambo Nickel SAS). As this slag cools down naturally under ambient conditions, its structure is fully crystallized. It is made of protoenstatite, clinoenstatite and forsterite. Its composition in wt.% of main metal oxides, measured by X-ray fluorescence, is SiO<sub>2</sub>: 52.1, MgO: 28.8 and Fe<sub>2</sub>O<sub>3</sub>: 14.4 (Augé and Touzé, 2015). It also exhibits traces of aluminium, manganese and calcium (Al<sub>2</sub>O<sub>3</sub>: 1.9%, CaO: 0.46%, MnO: 0.35%). This composition was found to be consistent with local analyses obtained from SEM-EDX elemental mapping of the particles (see **Supplementary Table S1**). As received samples were jaw

crushed, then ground in a dry ball mill to minus 100  $\mu$ m. The Particle Size Distribution (PSD) was measured by laser diffraction and yielded d<sub>10</sub>, d<sub>50</sub>, d<sub>90</sub>, and d<sub>32</sub> values of 7.5, 39.0, 100.5, and 12.7  $\mu$ m respectively (see **Figure 3**). Morphological analysis of 100,000 particles using an optical microscope (Morphologi G3, Malvern Instruments) gave an average HS (high sensitivity) circularity of 0.83 for their projected area (against 1.0 for a perfect circle). A closer look at the particles using both SEM and TEM images showed that particle surface exhibits some degree of roughness (see **Supplementary Figure S1**). This is in line with the fact that the specific surface area measured by nitrogen adsorption (1.25 m<sup>2</sup> g<sup>-1</sup>) is about ten times higher than that calculated from the PSD assuming smooth spherical particles (0.13 m<sup>2</sup> g<sup>-1</sup>).

A rotary sampler was used to produce equal mass (7.2 g) samples for carbonation testing. Based on Fe and Mg contents in the samples, the maximum amount of carbonate phases (FeCO<sub>3</sub> and MgCO<sub>3</sub>) is 0.0645 mole (or 5.85 g), and up to 0.0655 mole when including Ca and Mn. The other elements (Si and Al) do not form carbonate compounds.







# **Experimental Set-Up and Protocol**

The carbonation experiments were performed in a 300 ml selfmade stirred bead mill (**Figure 4**), built using an autoclave vessel suited to withstand pressure and temperature. Agitation was provided by a pin-type gas-inducing impeller. The mill tank was half-filled with millimetric grinding beads.

This investigation used either quartz sand (HN 0.6/1.6 from SIFRACO, 99% SiO<sub>2</sub>, 1.25–1.6 mm, 215 g) or AISI 420C stainless steel beads (1.6 mm in diameter, 675 g) as grinding media. Their density is 2.6 and 7.7 g cm<sup>-3</sup>, respectively, against 3.2 g cm<sup>-3</sup> for the slag. Quartz sand exhibits hardness of 7 on the Mohs' scale vs. 5–6 for enstatite.

The reactor was operated under controlled temperature and total pressure by means of a pressure regulator connected to a 200 ml CO<sub>2</sub> supply tank. The latter was instrumented with temperature ( $T_{\rm st}$ ) and pressure ( $P_{\rm CO2}$ ) gauges. The number of moles of CO<sub>2</sub> inside the tank was deduced from  $P_{\rm CO2}$  and  $T_{\rm st}$  values using Peng-Robinson's equation of state, which gave the instantaneous carbonation rate of the system. The duration of carbonation tests used in this work was 24 h. Once separated from grinding particles (on 800 µm sieve), filtered (on 0.22 µm filter) and oven dried at 70°C for 24 h, the carbonation products were analysed by two complementary techniques, namely

thermogravimetry (TGA) and elemental carbon content analysis. In the former analysis, carried out under nitrogen atmosphere, bound water was released during a 30 min plateau applied at 120°C. The weight loss between 300 and 600°C was then used to determine the carbonate amount (Bonfils et al., 2012), since the slag exhibits a constant mass in this temperature range. All of these measures (mass loss from TGA, total carbon content determination, total CO<sub>2</sub> consumption from storage tank) led to very consistent data regarding the final carbonation yield of the samples.

As part of a previous collaborative project (CNRT/ Carboscories), the carbonated slags were also characterized by electron microscopy (SEM/EDX), X-Ray Diffraction (XRD), and elemental analysis (ICP/AES analysis after acid digestion) by IMPMC and BRGM partners (Touzé et al., 2015).

Reference conditions inside the reactor were set to  $180^{\circ}$ C and 20 bar of CO<sub>2</sub> (leading to a total pressure of about 30 bar), a slurry concentration of 90 g of solid per liter of water and a stirring speed of 800 rpm.

In order to evaluate the benefit of the combined attritionleaching mode of operation, benchmark tests were conducted using a two-step carbonation mode. This consists in 24 h of attrition under ambient temperature and inert (azote) atmosphere, followed by 24 h of carbonation under the reference conditions without grinding beads.

# Thermodynamic and Kinetic Modelling of the Carbonation Process

This section lays the foundations for the thermodynamic and kinetic models used in this work to investigate the observed carbonation plateau observed in attrition-leaching carbonation experiments.

### Initial System Description

The description of the system was as follows: 1) the temperature *T* was considered constant and homogeneous within the reactor; 2) the amount of water was 80 g; 3) the amount of solid was 7.2 g; 4) an excess of CO<sub>2</sub> at temperature *T* and pressure  $P_{CO2}$  was available (accounting for the gas reservoir).

The slag was described as an assembly of oxides matching its elemental composition (see details in *Processed Material*):

- **System 1** included only the main three elements: Si, Mg and Fe, which accounts for 95.3 wt.% of the slag;
- **System 2** accounted also for the minor elements: Al, Ca and Mn, as they can contribute to the formation of silicate and/ or carbonate phases. For sake of illustration, Al is known, even as a trace element, to form a variety of species detrimental to carbonation, including alumino-silicates that can consume Mg at the expense of Mg-carbonates and Al-rich clay minerals instead of amorphous silica (Sissmann et al., 2014).

Under the conditions of the attrition-leaching carbonation process, the grinding beads themselves are potentially reactive and their chemical inertness was evaluated:

- a. Quartz sand was considered as an inert grinding medium, due to its high chemical stability (log  $K_{sp} = -2.37$  at 180°C) and low reactivity: its dissolution rate was reported to be  $10^{-15.3}$  mol cm<sup>-2</sup> s<sup>-1</sup> at 70°C from the acid to neutral pH range (Knauss and Wolery, 1988) against  $10^{-13.1}$  mol cm<sup>-2</sup> s<sup>-1</sup> for enstatite at 70°C and pH 5 (Oelkers and Schott, 2001).
- b. Stainless steel beads on the other hand are potentially corroded under the test conditions. They were accounted for by including a certain amount of  $(Fe_{0.87}Cr_{0.13})$  in the initial materials description (see *Effect of the Nature of the Grinding Beads on the Carbonation Products*).

### Thermodynamic Model

Given the chemical constitution of the system, the thermodynamic model needs to account for multiphasic equilibria including a pressurized gaseous phase, an aqueous electrolytic phase and various solid compounds and their mixtures (i.e., solid solutions).

Equilibrium calculations were carried out using PHREEQC, a simulator designed to perform a wide variety of aqueous geochemical calculations, including high-pressure conditions (Parkhurst, 1995; Parkhurst and Appelo, 1999; Zhu and

Anderson, 2002; Parkhurst and Appelo, 2013). In relation to the system constitution, this software implements the Peng-Robinson equation of state to describe gaseous mixtures at high pressure, three activity coefficient models to describe the aqueous phase (namely extended Debye-Hückel [the so-called B-dot equation, Helgeson (1969)], Specific Ion Theory (Biederman, 1986) and Pitzer (Pitzer, 1973), as well as solid solutions with binary interaction parameters (Glynn and Reardon, 1990).

In the present system, the very small ionic strength of the aqueous solutions (about  $10^{-3}-10^{-2}$  mol/kg<sub>water</sub>) justified the use of the B-dot equation for activity coefficient calculations. Accordingly, this work used the Thermoddem 2017 database due to its ability to satisfactorily predict the solubility of possible slag carbonation species, including magnesium carbonates, iron carbonates (Fosbøl et al., 2010; Bénézeth et al., 2011) and a wide variety of alumino-silicate phases (Blanc et al., 2012; Blanc, 2017). This database selection is different from our previous work (Cassayre et al., 2016), which implemented the LLNL database. The decision of switching to the Thermoddem 2017 database is mostly motivated by an improved assessment of the SiO<sub>2(aq)</sub> specie properties, which takes into accounts more recent experimental data and evaluation from Rimstidt (1997) and Gunnarsson and Arnórsson (2000).

As for  $CO_2$ -water vapour mixture, the simulations used Peng-Robinson equation of state to describe fugacities and thus to calculate the  $CO_2$  solubility in water according to Henry's law.

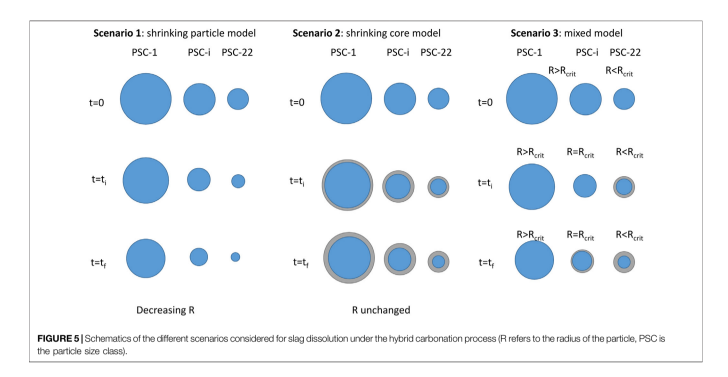
Regarding solids, all mineral phases present in the database that include these elements were allowed to precipitate, with the exception of quartz due to the existence of a strong kinetic barrier that favours the formation of amorphous silica (Daval et al., 2011). Direct measurements confirmed this hypothesis. Indeed, with the exception of the experiments using sand as grinding media, XRD, and EDX analyses never detected quartz in the solid product, but an increased amount of Si-rich amorphous phase(s).

Precipitated carbonates have almost identical crystal structures (Effenberger et al., 1981) and can form substitutional solid solutions on a large composition range. Solid solutions are more stable than the separate phases and commonly exhibit deviations from ideal behaviour, hence the need to describe their thermodynamic properties accurately. However, PHREEQC only allows the definition of excess free energy for binary solid solutions. Therefore a (Mg,Fe)CO<sub>3</sub> solid solution with a regular solution parameter of 4.44 kJ/mol (Chai and Navrostky, 1996) was considered for system 1 and an ideal (Mg, Fe, Ca, Mn) CO<sub>3</sub> solid solution for system 2.

### Kinetic Modelling of the Slag Dissolution

Slag dissolution (see **Figure 1**) is here considered as the ratelimiting step, owing to the low acidity provided by  $CO_2$  hydrolysis (the predicted pH of the system – including the basic slag – being about 5 at equilibrium). Consequently, both dissolved gas and formed solid products were supposed to be at thermodynamic equilibrium at each time step during the dissolution process.

To describe the slag leaching kinetics, three distinct scenarios (**Figure 5**) were envisaged in this paper. Although they do not account for the actual elementary comminution processes, they



consider the main reaction mechanisms that govern carbonation yield under attrition conditions. These scenarios depend on the concomitance between attrition and carbonation and the presumed efficiency of attrition for surface layer removal:

(1) Scenario 1: A *shrinking particle mechanism* is assumed to represent the attrition-leaching process. In this case, attrition peels off completely (and only) the surface layers as they form on the reactive particles. Surface reaction is the rate-determining step in scenario 1 and the corresponding mass balance is represented by **Eq. 1** for a given particle size class:

$$\frac{dn_i}{dt} = -r \cdot SSA_{i0} \cdot m_{i0} \left(\frac{n_i(t)}{n_{i0}}\right)^{2/3}$$
(1)

where  $n_{i0}$  and  $n_i$  are the initial and current number of moles of slag in particle size class i,  $m_{i0}$  the corresponding initial weight of slag (kg),  $SSA_{i0}$  the initial specific area (m<sup>2</sup> kg<sup>-1</sup>) of the particle size class i.

The exponent 2/3 in **Eq. 1** assumes an idealized spherical shape of the particles. For consistency, calculation of  $SSA_{i0}$  used the surface-to-volume ratio of the smooth sphere whose diameter is that of particle size class *i*. The consequence of this assumption is discussed in *Scenario 1: Shrinking Particle Model*. Note that the attrition process is expected to modify the surface aspect of the slag particles, which means that their actual shape factor is likely to vary in time.

The surface reaction rate, r, is expressed by:

$$r = k_0 \cdot exp\left(\frac{-E_A}{\mathbf{R} \cdot \mathbf{T}}\right) a^n_{(H^+)} a^m_{(Mg^{2+})} \left(1 - 10^{SI}\right)$$
(2)

Given the mineralogy of the slag, the saturation index *SI* and kinetic parameters ( $k_0$ ,  $E_A$ , n, m) are those of enstatite, which are estimated from the work of Oelkers and Schott (2001):  $k_0 = 2.4$  mol m<sup>-2</sup> s<sup>-1</sup>,  $E_A = 48.5$  kJ mol<sup>-1</sup>, n = 1/4 and m = -1/8 (*a* being the activity of the ion considered).

(2) Scenario 2: Slag dissolution mechanism obeys a *shrinking core mechanism*, while product layers form on the surface of the unreacted slag core. Diffusion within the solid layers is the rate-determining step for this scenario. This applies to the conventional aqueous carbonation process without attrition, for which the formation of surface layers has been already reported (Schott and Berner, 1983; Daval et al., 2011; Julcour et al., 2015). An illustration can be found in **Supplementary Figure S2**. Therefore, the two-step experiment, where attrition and carbonation are sequential (cf. *Experimental Set-Up and Protocol*), can be used to evaluate the corresponding diffusion parameter.

The associated mass balance is described by Eq. 3, derived from Levenspiel (1998), for a given particle size class:

$$\frac{dn_i}{dt} = -\nu_{S/H^+} \frac{D_{eH^+} \cdot C_{H^+ext}}{R_{i0}} \cdot SSA_{i0} \cdot m_{i0} \frac{\left(\frac{n_i(t)}{n_0}\right)^{1/3}}{1 - \left(\frac{n_i(t)}{n_0}\right)^{1/3}}$$
(3)

where  $n_{i0}$  and  $n_i$  are the initial and current number of moles of slag in particle size class i,  $m_{i0}$  the corresponding initial weight of slag (kg),  $SSA_{i0}$  the initial specific area (m<sup>2</sup> kg<sup>-1</sup>) of the particles of mean radius  $R_{i0}$  (equal to that of the sphere of the same radius),

 $D_{eH^+}$  and are respectively the effective diffusivity and external (bulk) concentration of the proton,  $\gamma_{S/H^+}$  the ratio of the

stoichiometric coefficient of slag to that of proton in the dissolution equation (0.5).

Note that **Eq. 3** is not defined at time zero of the dissolution when  $n_i = n_{i0}$ . Therefore a limitation by the surface chemical reaction (see **Eq. 1**) is first supposed until the conversion of slag  $\left(1 - \frac{n_i(t)}{n_{i0}}\right)$  reaches 0.05 to prevent any numerical stiffness (values between 0.01 and 0.1 showing no significant effect on the results).

(3) Scenario 3: This scenario mixes scenarios 1 and 2. Indeed, it considers that attrition is no longer effective for surface layer removal below a critical slag particle diameter. In other words, scenario 3 means that slag particles larger than the critical diameter undergo scenario 1 while particles finer than the critical size are subject to scenario 2. It also means that particles with an initial diameter larger than the critical size might experience both scenarios during the attrition-leaching carbonation process.

Scenarios 1 to 3 are limiting cases. Several other scenarios, the likelihood of which cannot be denied will be considered in the future. One of them corresponds to attrition events peeling off more than the passivating surface layers, and producing fine unreacted slag particles at the same time. While this scenario is expected to yield an increased carbonation rate compared to scenario 1, it is also likely to be more energy intensive. Indeed, the least energy intensive attrition-leaching carbonation conditions are those that only use the energy necessary to exfoliate surface layers, hence the value of scenario 1. Fragmentation events, also likely to occur during attritionleaching carbonation, are not considered either in the three selected scenarios.

All these models require the description of the particle size distribution (PSD) of the feed material. It was modelled using a Rosin-Rammler distribution, and then discretized into 22 size classes. Individual size classes were entered into the geochemical code as a distinct material, with the same thermodynamic properties but different specific surface area ( $SSA_{i0}$ ) and amount ( $m_{i0}$ ). The shrinking core model used the PSD measured after the 24 h attrition stage of the two-step process in order to evaluate the effect of size reduction possibly occurring as initial step of the attrition-leaching process in scenario 2.

Figure 3 shows the corresponding cumulative PSDs and reveals that the  $d_{50}$  shifts from 39.0 to 8.1 and 2.8 µm after 24 h of attrition using stainless steel beads and quartz sand, respectively. The strongest reduction of the mean particle size observed in the latter case is explained by the self-attrition of the quartz sand particles, resulting into an increase in weight of the final solid product much higher than the CO<sub>2</sub> uptake. Nevertheless, a different dominant comminution mechanism (breakage for steel beads vs. attrition for quartz sand) cannot be excluded to explain such a difference in the resulting PSDs, due to the high density difference between these grinding particles.

# **RESULTS AND DISCUSSION**

Thermodynamic considerations on the nature of the precipitated phases are discussed first. Then, the different mechanisms considered to describe the slag dissolution kinetics are assessed against the carbonation rates measured during the two-step and concomitant attrition-leaching processes.

# Thermodynamic Predictions of Slag Dissolution Under Attrition-Leaching Carbonation Process Conditions

# Product Speciation and Effect of Minor Elements on the Operating Window of Slag Carbonation

**Figure 6** shows equilibrium calculations for the system  $SiO_2$ -MgO-Fe<sub>2</sub>O<sub>3</sub>-CO<sub>2</sub>-H<sub>2</sub>O (system 1), which accounts for the major elements in the slag. **Figure 6A** gives the carbonation yield as a function of temperature and CO<sub>2</sub> partial pressure, while **Figure 6B** shows the nature and amount of solid products at 20 bar of CO<sub>2</sub>.

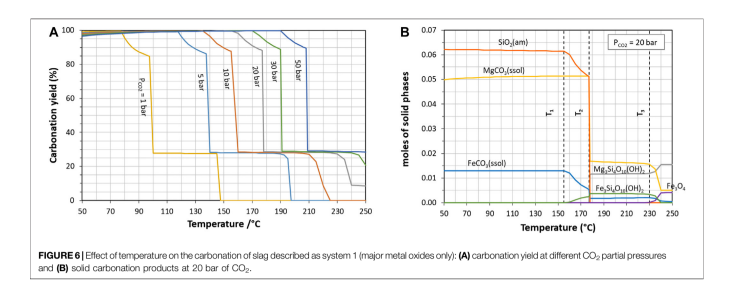
At a given CO<sub>2</sub> partial pressure, mineral speciation varies with temperature in the range investigated (50–250°C), which results in a sharp decrease in carbonation extent beyond a certain temperature. In **Figure 6B**,  $T_1$  delineates the most suitable operating window for quantitative carbonate yield, in which leached silicon precipitates as amorphous silica, while leached iron and magnesium precipitate as a carbonate solid solution (ssol). At higher temperatures ( $T > T_1$ ), ferrous talc (minnesotaite), then magnesium talc ( $T > T_2$ ) and magnetite ( $T > T_3$ ) form at the expense of carbonates. Owing to the composition of the slag (with Mg >> Fe), the strongest reduction in carbonation extent coincides with  $T_2$ . It should be noted that the reference conditions under investigation (180°C, 20 bar of CO<sub>2</sub>) stand beyond this second limit, while measured carbonation yield actually exceeded 50%.

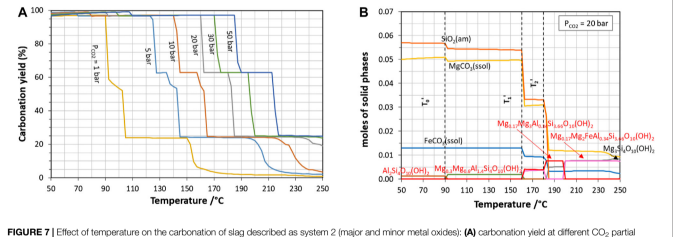
The same pattern is found across the whole  $CO_2$  pressure range, with only a shift for the threshold temperatures, and the occurrence of antigorite for  $P_{CO2}$  less than 20 bar.

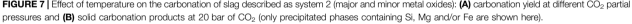
Our recent paper (Bourgeois et al., 2020) confirmed the existence of such pivot conditions for the carbonation of a water-quenched nickel slag, prompting a strong reduction of the extent of  $CO_2$  fixation (from 75% down to 25%) when further increasing temperature or reducing  $P_{CO2}$ . The exact values of these thresholds are very sensitive to the equilibrium constants of the predicted phases, as shown below.

Including the minor elements of the slag (system 2), carbonation yield profiles (**Figure 7A**) exhibit not only one, but two successive steep drops (at  $T'_1$  and  $T'_2$ ), resulting for instance in a maximum yield of about 60% just below 180°C at 20 bar of CO<sub>2</sub>. This is due to the precipitation of various stable alumino-silicates that consume Mg and/or Fe (**Figure 7B**): montmorillonite (Mg<sub>0.3</sub>Mg<sub>0.6</sub>Al<sub>1.4</sub>Si<sub>4</sub>O<sub>10</sub>(OH)<sub>2</sub>) and saponite-like phases (Mg<sub>0.17</sub>Mg<sub>3</sub>Al<sub>0.34</sub>Si<sub>3.66</sub>O<sub>10</sub>(OH)<sub>2</sub>).

Measured carbonation yield as high as 50% (corresponding to 0.032 mole of carbonates in the reactor) were obtained after 24 h  $\,$ 







of carbonation under the reference conditions, using quartz sand as grinding medium. X-ray diffraction analysis of the solid product showed initial mineral phases, quartz from sand grinding particles and mixed crystallized carbonates. Further examination with a transmission electron microscope pointed out that the solid product was made of an intimate mixture of nano-sized carbonates and silicate grains, the latter having different contents of magnesium, iron and aluminium and various structures (globules or fluffy aggregates) (see **Figure 8A** in *Effect of the Nature of the Grinding Beads on the Carbonation Products*). Although the exact nature of the remaining alumino-silicate phases has not been fully identified, these observations are consistent with the thermodynamic simulations, which predict the formation of a variety of alumino-silicate phases.

# Effect of the Nature of the Grinding Beads on the Carbonation Products

To further test the reliability of the thermodynamic predictions, the case of stainless steel grinding beads was examined closely given their expected reactivity. Under the carbonation condition, a significant increase in  $CO_2$  consumption was measured, reaching up to 0.045 mole in 24 h. Such a value corresponds to a theoretical 70% carbonation yield. Carbonate amounts calculated from elemental C analysis were systematically higher - by as much as 15% - than those obtained from TGA analyses. This discrepancy, not observed in the previous experiments, raised the suspicion of a possible product contamination. Moreover, XRD revealed an enhanced formation of siderite (FeCO<sub>3</sub>) and mixed (Mg,Fe)CO<sub>3</sub> carbonates, while ICP/AES

analysis after acid digestion revealed that the iron to magnesium ratio jumped from 0.25 to 0.45 between the initial material and the final solid products.

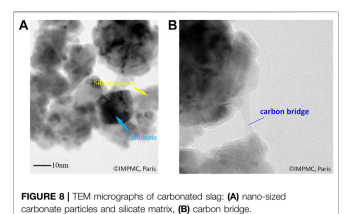
In order to elucidate this chemical influence of stainless steel balls, the amount of iron-chromium solid solution was increased in the initial simulated system, up to the increment of iron measured in the product of about 0.01 mole (corresponding to 0.1% of the beads). This stainless steel corrosion behaviour under low solution acidity was reported in a previous contribution (Cassayre et al., 2016), which used a different thermodynamic database (see *Thermodynamic Model*) and led to somewhat different results for iron speciation since several (alumino) ferro-silicates were missing in the LLNL database.

Thermodynamic simulations performed at 175°C (just below  $T_2$  or  $T'_2$ ) predicted that the iron extracted from the steel beads could be converted to either minnesotaite or siderite and Fe-Mg saponite (with an increase of magnesite), depending on the system considered (Figure 9). These results prove that an accurate thermodynamic description of the various phyllosilicates formed is also of primary importance to evaluate the contribution of iron pollution to the carbonation yield. With the reference conditions, the simulation predicted that 0.048-0.057 mole of carbonates could be formed at the most when taking into account the iron dissolved from beads, but only the most complete system (system 2) could predict its transformation into siderite.

The simulation also predicts the formation of solid carbon as a by-product of iron oxidation, according to the following siderite forming reaction:

$$Fe_{(steel)} + 3/2CO_{2(aq)} = FeCO_{3(ssol)} + 1/2C_{(s)}$$
 (4)

TEM micrographs confirmed the presence of carbon bridges between particles (**Figure 8B**), which supports the formation of solid carbon. Furthermore, the occurrence of this reaction is consistent with the higher carbon content measured by carbon analysis (carried out in oxidative conditions, and thus including elemental C) compared to the carbonates amounts determined by TGA (carried out in nitrogen atmosphere and thus not including elemental C). This observation also demonstrates the interest of

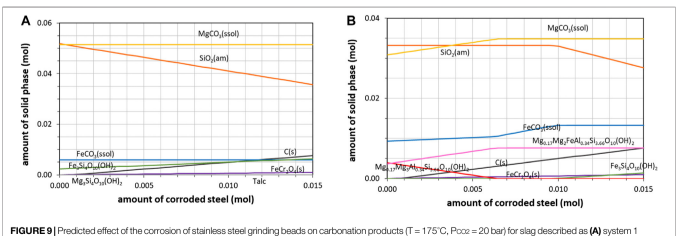


combining both analytical techniques in carbonation products analysis.

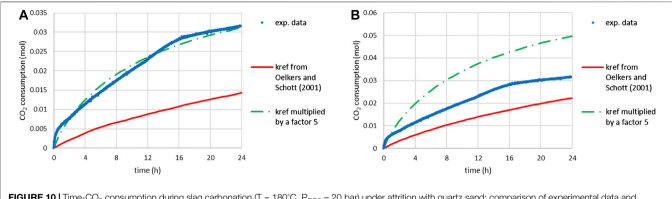
# Kinetic Modelling of Slag Dissolution During the Attrition-Leaching Carbonation Process

Operating particle leaching in a stirred bead mill leads to a change in PSD due to several concurrent processes: dissolution of the reactive slag, total or partial abrasion of surface product layers, the effectiveness of which may vary over time mainly due to its particle size dependency, and particle breakage. Instead of modelling these coupled phenomena, which is out of reach at this stage of our work, this section has no other objective than trying to identify the mechanisms that contribute the most to the carbonation extent. To this end, three limiting cases defined in *Kinetic Modelling of the Slag Dissolution* as scenarios 1 to 3 are used. They are briefly recalled hereafter:

- Scenario 1 is a purely surface-controlled (or reactioncontrolled) process, which corresponds to an effective attrition process that peels off passivation layers as they form. It is described by the shrinking particle model;
- Scenario 2 is a purely diffusion-controlled process, which corresponds to a conventional aqueous carbonation process



(major metal oxides only) and (B) system 2 (major and minor metal oxides).



**FIGURE 10** | Time-CO<sub>2</sub> consumption during slag carbonation ( $T = 180^{\circ}C$ ,  $P_{CO2} = 20$  bar) under attrition with quartz sand: comparison of experimental data and predictions of scenario 1 (assuming a reaction-limiting process): (**A**) for system 2 (including main and secondary elements) and (**B**) for system 1 (limited to the main elements) with inhibition of minnesotaite precipitation.

with surface passivation. It is described by the shrinking core model. It is also hypothesized that the change in particle size distribution by the grinding occurs much faster than the slag dissolution, so that the shrinking core model uses the PSD after attrition stage;

• Scenario 3 is a mixed process. It corresponds to a reactioncontrolled process (scenario 1) for particles larger than a critical size, above which attrition is able to continuously remove surface layers. For particles finer than the critical size, this scenario switches to a diffusion-controlled process (scenario 2).

In the view of an efficient mineral carbonation, scenario 1 is considered the most favourable among the three, since it is not limited by any diffusion barrier. Furthermore, as aforementioned, attrition of the remaining reactive core (beyond the peeling off of the product surface layer) or particle fragmentation are not discussed here.

### Scenario 1: Shrinking Particle Model

**Figure 10** compares the carbonation rates measured with quartz sand to those calculated with this scenario, thus assuming that reaction on fresh surfaces is the rate-controlling step. For the sake of comparison with the experimental data, the simulations were performed at 175°C, that is thus below temperature  $T_2$  or  $T_2$ .

Scenario 1 was first applied to system 2 (where the slag is described as its major and minor metal oxides), using the feed material PSD and the reaction rate law established by Oelkers and Schott (2001) for enstatite. Although the upper bound of the temperature range (28–168°C) over which Oelkers and Schott fitted their reaction rate is close to our experimental conditions (180°C), **Figure 10A** shows that scenario 1 largely underestimates the measured carbonation kinetics. It was necessary to increase the kinetic rate constant ( $k_{ref}$ ) by a factor 5 to satisfactorily reproduce the experimental carbonation kinetics. This result is very likely influenced by the underestimation of the actual surface area of the particles from the measured PSD and the smooth spherical particle assumption. As mentioned in *Processed Material*, the ratio between the two surface areas was close to 10 for the initial slag particles.

After adjustment of the rate constant, the model predicts that conversion of the slag particles would have reached 77% in 24 h and the formation of alumino-silicate by-products containing Mg and/or Fe would explain the lower carbonation yield (of about 50%).

Scenario 1 was then applied to system 1 (where the slag is described as its major metal oxides only). Minnesotaite was excluded from the possible carbonation products as per the conclusions drawn in *Effect of the Nature of the Grinding Beads on the Carbonation Products*, and possible carbonation products were limited to the desired ones, namely siderite, magnesite and amorphous silica. It is noted that these phases were all that was observed to form by Bourgeois et al. (2020) during the attrition-leaching carbonation of another nickel slag. **Figure 10B** shows that scenario 1 predicted the very first moments of the reaction, but yielded too high a CO<sub>2</sub> uptake, since in this case the carbonation yield always matches the conversion of slag particles (no Mg/Fe silicate by-product).

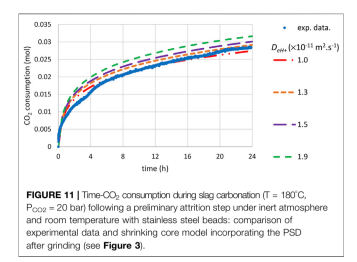
Thorough examination of XRD pattern of the product gave clues on the actual conversion of the slag: Rietveld refinement indicated that the remaining slag phases (forsterite, enstatite, augite, diopside) accounted for up to 50 wt.% of the crystalline Mg containing phases (and carbonates the rest) (Ploquin et al., 2016).

Therefore, the "fitted" shrinking particle model seems in fact to overestimate the slag conversion and the potential phyllosilicate by-products are most likely different from those predicted by the thermodynamic calculations, as mainly in the form of amorphous phases.

For the examination of scenarios 2 and 3, the simulated material was thus reduced to system 1, with inhibition of minnesotaite precipitation.

### Scenario 2: Shrinking Core Model

This model (see **Eq. 3**) requires the value of the effective diffusion of the proton in the surface leach layer,  $D_{eH^+}$ . This value was first estimated based on the carbonation rates measured during the two-step carbonation process, before scenario 2 was assessed with respect to the results of the attrition-leaching carbonation process using quartz sand.



# Estimation of the Effective Diffusivity $D_{eH^+}$ in the Product Surface Layer

In the two-step mode, the slurry bearing the slag particles is first subjected to attrition under inert atmosphere and room temperature. The slurry and grinding beads are then separated by screening, after which the slurry undergoes a regular carbonation step, during which the surface of the slag particles is progressively covered by a product layer. Thus, knowing the PSD after attrition stage, the shrinking core model can be tuned on the corresponding experimental  $CO_2$  uptake.

To avoid any misevaluation of the product PSD after attrition due the presence of debris from the grinding medium, the model was applied to the case of stainless steel beads to estimate  $D_{eH^+}$ and used the corresponding PSD shown in **Figure 3**. This supposed that the product layer is essentially the same regardless of the initial attrition step (using stainless steel beads or quartz sand).

**Figure 11** shows that a satisfactory agreement between simulated and measured carbonation rates was found for a fitted value of  $D_{eH^+}$  equal to  $1-1.3 \times 10^{-11} \text{ m}^2 \text{ s}^{-1}$ . This value lies inside the range of values reported in the literature for the diffusivity of proton in product layers formed during acid digestion or aqueous carbonation of magnesium of calcium silicates under various conditions – ranging from  $10^{-16}$  to  $5.10^{-10} \text{ m}^2 \text{ s}^{-1}$  (Daval et al., 2009; Van Essendelft and Schobert, 2009; Bonfils, 2012). Such a wide range can be ascribed to the variety of the product layers formed, from nanometre-thick and totally impervious at low temperature (of about  $100^{\circ}$ C) to micrometre-thick and fibrous-like at higher one (Julcour et al., 2014).

The shrinking core model assumes that the fluid reactant migrates to the core surface through the porous structure of the surface layer. The effective diffusivity is then linked to the molecular diffusivity  $D_{mH^+}$  according to:

$$D_{eH^+} = \frac{\varepsilon_p}{\tau} D_{mH^+} \tag{5}$$

where  $\varepsilon_p$  and  $\tau$  refer to the porosity and pore tortuosity of the product layer.

The molecular diffusion coefficient of proton in water at 180°C can be estimated to  $3.9 \times 10^{-8}$  m<sup>2</sup> s<sup>-1</sup> according to the correlation of Lee and Rasaiah (2011). Considering a tortuosity value of about five for low-porous matrices (Ghanbarian et al., 2013), this yields a surface layer porosity of the order of 0.1%. This is not unrealistic according to the conclusions of Van Essendelft and Schobert, who accept up to 3 orders of magnitude difference between the two diffusivities.

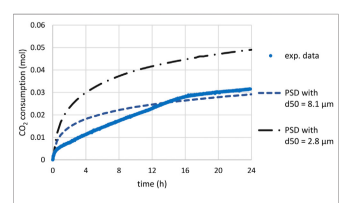
### Assessment of Scenario 2 for the Attrition-Leaching Carbonation Process

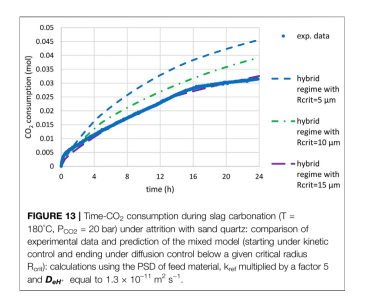
The shrinking core model using the estimated  $D_{eH^+}$  value was then applied to the attrition-leaching carbonation process with quartz sand. Assuming that shift in PSD shown in **Figure 3** occurs very fast, scenario 2 was applied to this PSD. Keeping in mind that fines produced by attrition of quartz sand itself have probably contributed in part to the larger change noted in **Figure 3**, both post-attrition PSDs were considered here. As shown in **Figure 12**, scenario 2 yielded the overall trend and magnitude of the carbonation kinetics obtained for the coarser PSD. However, it predicted a too early slowing down of the measured kinetics.

Examination of scenarios 1 and 2 thus led to the conclusion that a mixed mechanism in between the two could have occurred to explain the experimental trend.

#### Scenario 3: Mixed Model

As discussed in *Scenario 1: Shrinking Particle Model*, the formation of undesirable silicate phases can lower the equilibrium carbonation yield. The slowing down of the attrition-leaching carbonation kinetics could also be ascribed to a gradual decrease in attrition efficiency related to the increasing fineness of the reactive particles. Several factors contribute to this statement. While attrition may increase the reactivity of reactive surfaces by amorphization, finer particles exhibit less surface defects and require more energy for their surface fragmentation (Kwade and Schwedes, 2007; Schilde et al., 2010). The finer the particles, the lesser the likelihood of particles to experience shear stress events inside the stirred bead mill.





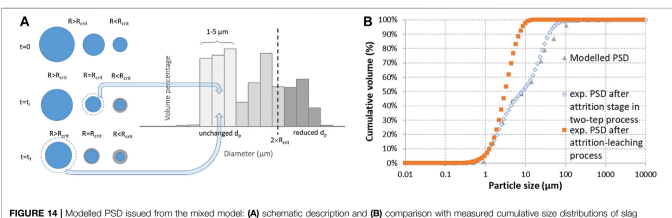
Although this may not apply here as tested slurries were dilute (~3 vol.%), increased slurry fineness may yield a rise in slurry viscosity that reduces attrition efficiency. Scenario 3 proposes a mixed regime that could possibly approximate the drop in efficiency of attrition with increased fineness. This scenario subjects particles larger than a critical size to a reaction-controlled process (scenario 1), and finer particles to a surface-controlled process (scenario 2). **Figure 13** shows the simulated prediction for scenario 3, for three critical particle sizes, using the modified kinetic rate constant (multiplied by a factor 5) and the value of  $D_{eH^+}$  previously optimised.

The third limiting scenario matches the experimental trend for a critical radius  $R_{crit}$  of 15 µm. This would mean that feed particles initially smaller than 30 µm in diameter would not change in size (owing to the shrinking core mechanism) while coarser particles would be reduced down to this diameter value. Attrition fragment would be much finer though (in the micrometer size range).

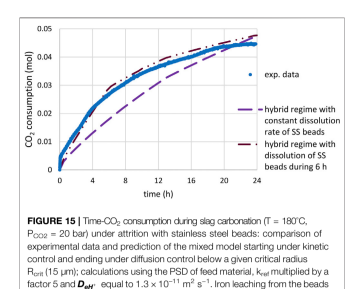
Based on the calculated conversion of the different size classes and assuming a similar molar volume of the reactant and the product mixture, this would yield 38% of the initial volume transformed into small fragments. The rest would correspond to a volume percentage of 13% for particles above 100 µm in diameter, 16% in between 30 and 50 µm, and 33% for those unchanged in size (below 30 µm in diameter in the initial PSD). As abovementioned, the particle size measured after attrition with quartz sand was affected by the autogenous attrition of quartz sand particles themselves, quartz contributing up to 60 wt.% of the crystalline phases for the hybrid process (Ploquin et al., 2016). With stainless steel beads, part of the product came from the dissolution of iron from the beads. Therefore, the PSD measured after attrition alone with stainless steel beads could be viewed as a better approximation of the final product's PSD. As illustrated in Figure 14, the mixed scenario may be plausible when compared to this experimental size distribution (the smallest fragments being distributed equally over the 1-5 µm size classes in the modelled PSD). It should be noted, however, that the maximum fineness  $(d_{50})$  achievable in industrial stirred bead mills is down to 1/1,000 of the grinding medium size, but in practice it could be in the order of 200-500 times smaller than the bead size used (Hassal, 2008). This would correspond here to 2-10 µm particles, since the experiments used 1.6 mm grinding medium.

Finally, in **Figure 15**, the predictions of scenario 3 (with  $k_{ref}$  multiplied by 5,  $R_{crit} = 15 \,\mu\text{m}$  and  $D_{eH^+} = 1.3 \times 10^{-11} \,\text{m}^2 \,\text{s}^{-1}$ ) are compared to the experimental time-CO<sub>2</sub> consumption data obtained during attrition-leaching carbonation with stainless steel beads. Assuming a constant leaching rate of the iron from steel (up to the final amount measured), the model would only match the end point. To follow the whole curve, this corrosion process should have mainly occurred during the first 6 h, which would mean that a passivation layer was also progressively formed on the beads and not fully peeled off.

Although based on a simplified mechanism, explaining the evolution of the material PSD through the attrition of product layer in small fragments as long as the slag particles remain large enough, scenario 3 could conciliate the measured carbonation rate, the final slag conversion and the final particle size.



particles after 24 h of attrition grinding using stainless steel beads (in two-step and attrition-leaching processes) (conditions of Figure 13).



# **CONCLUSIONS AND PERSPECTIVES**

is assumed to occur throughout the reaction or during the first 6 h only.

Owing to the continuous renewal of reactive surfaces that goes beyond a simple particle size reduction effect, the attritionleaching carbonation process has been shown to strongly improve the carbonation yield of natural minerals and industrial residues alike. Invariably, the attrition-leaching carbonation yield levels off 20-50% below the yield expected from stoichiometry with all feedstocks tested to date, including the nickel slags used in this study. Using an original thermokinetic modelling methodology that combines equilibrium thermodynamic and particle reaction kinetic models, this paper attempts to shed light on the mechanisms that govern this behaviour. Indeed, it is thought that this repeatable behaviour can originate either from the precipitation of undesirable silicate by-products whose formation competes with that of carbonates under single-step aqueous mineral carbonation conditions, or from a kinetic limitation specific to the attrition-leaching carbonation process itself.

Thermodynamic modelling is a powerful tool that can predict the likelihood of the formation of precipitated species, accounting for all the inputs to the process. In particular, this work has shown that it is invaluable for anticipating the possible effects of slag impurities and grinding medium on the aqueous carbonation system. The work also confirms that thermodynamic modelling cannot be used in isolation, as it must be accompanied by thorough analysis of the products by various complementary physico-chemical techniques for quantitative validation and use. The results obtained by thermodynamic modelling for the systems selected in this study, namely nickel slag with either stainless steel or quartz sand grinding media, show that the observed slowing down of the carbonation process could be explained by the formation of Mg- and/or Fe-rich silicates that precipitate at the expense of carbonates.

While the conclusion is not yet clear-cut, this work shows that a more likely explanation is of a kinetic nature. The kinetic limitation in question corresponds to a gradual reduction in attrition effectiveness as reactive feed particles leach out. Indeed, while neither the shrinking particle nor the shrinking core model could match the slowing down of the carbonation process, a mixed model that combines the two reproduced the complex course of the process satisfactorily. This model was built using a separate evaluation of the underlying parameters, in particular the surface reaction rate constant and diffusivity of the leached layer, as well as the measured evolution of the particle size distribution.

Pending additional confirmation, this finding is positive for the improvement of the attrition-leaching carbonation process. Indeed, if the carbonation limitation were of a thermodynamic nature, nothing simple could be done to improve carbonation yield of the attrition-leaching carbonation process. However, the likelihood of the limitation being of a kinetic nature opens the possibility to increase the carbonation yield by improving the performance of the attrition process itself. This requires however that the workings of the attrition process be understood at a local (particle) level in the context of the attrition-leaching carbonation process. As suggested by the good results obtained with the mixed model, the intrinsic limit of the attrition process to grind fine particles beyond a critical size is probably an important feature of the problem, along with the size-dependent efficiency of the attrition process. Notwithstanding their size, the choice of grinding media is also a critical component of the problem, as Ni slag for instance is a particularly abrasive carbonatable feedstock. Another parameter of importance is the rheology of the system, which was uncontrolled in the batch experiments used in this study. Other complex factors may be at work here also, considering that the attrition operates on particle surfaces whose mineralogical, mechanical and reactive properties are affected locally by mechanical stresses. Understanding these factors is inseparable from conducting extensive characterisation of the phases present - composition, nature and quantity - at different reaction times, which is most challenging.

While it is believed that the optimization of the attritionleaching carbonation process lies in finding the right equilibrium between attrition and leaching rates, a comprehensive investigation of the attrition mechanisms at the particle level is now mandatory to elucidate and optimize the inner workings of the attrition-leaching carbonation process.

# DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

# **AUTHOR CONTRIBUTIONS**

CJ and FB wrote the manuscript. The attrition-leaching carbonation process is an original process proposed by FB. LC developed the theoretical framework of the thermodynamic modelling and provided input and critical feedback to the manuscript. CJ devised the main conceptual ideas of the kinetic modelling and performed the numerical calculations.

IB carried out the experiments and JD contributed to the modelling work, under the supervision of FB, CJ, and LC. All authors analysed the data and discussed the results.

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# REFERENCES

- Augé, T., and Touzé, S. (2015). Caractérisation des résidus de pyrométallurgie de l'usine de Koniambo. Rapport BRGM/RC-64842-FR.
- Bénézeth, P., Saldi, G. D., Dandurand, J.-L., and Schott, J. (2011). Experimental determination of the solubility product of magnesite at 50 to 200°C. *Chem. Geol.* 286, 21–31. doi:10.1016/j.chemgeo.2011.04.016
- Biederman, G. (1986). "Introduction to the specific interaction theory with emphasis on chemical equilibria," in *Metal complexes in solution*. Editors E. A. Jenne, E. Rizzarelli, V. Romano, and S. Sammartano (Padua, Italy: Piccin), 303–314.
- Blanc, P. (2017). Thermoddem: update for the 2017 version. Report BRGM/RP-66811-FR. Available at: https://thermoddem.brgm.fr/sites/default/files/upload/ documents/brgmrp-66811-fr\_final\_report.pdf (Accessed November 3, 2020).
- Blanc, P., Lassin, A., Piantone, P., Azaroual, M., Jacquemet, N., Fabbri, A., et al. (2012). Thermoddem: a geochemical database focused on low temperature water/rock interactions and waste materials. *Appl. Geochem.* 27 (10), 2107–2116. doi:10.1016/j.apgeochem.2012.06.002
- Bodor, M., Santos, R. M., Van Gerven, T., and Vlad, M. (2013). Recent developments and perspectives on the treatment of industrial wastes by mineral carbonation - a review. *Cent. Eur. J. Eng.* 3 (4), 566–584. doi:10. 2478/s13531-013-0115-8
- Bonfils, B. (2012). Mécanismes et verrous de la carbonatation minérale du CO2 en voie aqueuse. PhD thesis. Toulouse (France): Institut National Polytechnique de Toulouse. Available at: https://oatao.univ-toulouse.fr/7920/1/bonfils\_partie\_1\_sur\_2.pdf.
- Bonfils, B., Julcour-Lebigue, C., Guyot, F., and Bodénan, F., Chiquet, P., and Bourgeois, F. 2012). Comprehensive analysis of direct aqueous mineral carbonation using dissolution enhancing organic additives. *Int. J. Greenh. Gas Control.* 9, 334–346. doi:10.1016/j.ijggc.2012.05.009
- Bourgeois, F., Laniesse, P., Cyr, M., and Julcour, C. (2020). Definition and exploration of the integrated CO<sub>2</sub> mineralization technological cycle. *Front. Energy Res.* 8, 1-17. doi:10.3389/fenrg.2020.00113
- Cassayre, L., Bourgeois, F., Julcour-Lebigue, C., Benhamed, I., Diouani, J., and Nahdi, K. (2016). "Defining the operating conditions of the attrition-leaching process using thermodynamic process modelling," in IMPC 2016: XXVIII international mineral processing congress proceedings, Québec, Canada, September 2016, 2490–2501.

with expertise in geochemistry, mineralogy, crystallography (Institut de Minéralogie, de Physique des Matériaux et de Cosmochimie, Paris and Bureau de Recherches Géologiques et Minières, Orléans), chemical engineering (Laboratoire de Génie Chimique, Toulouse) and industrial integration (BRGM). The authors wish to thank Solène Touzé (BRGM) for the coordination of the project and ICP/AES analyses of the products, Thierry Augé (BRGM) for the characterization of the KNS slag, Florian Ploquin and Farid Juillot (IMPMC) for the DRX/Rietveld analyses and François Guyot (IMPMC) for the TEM images of the products. They also acknowledge the technical team of LGC, Jean-Louis Labat and Lahcen Farhi for their support on the experimental set-up and Christine Rey-Rouch for her help on thermogravimetry, particle morphology and size analyses, as well as Isabelle Borget (LCC Toulouse) for CHN measurements.

# SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fceng.2020.588579/full#supplementary-material

- Chai, L., and Navrotsky, A. (1996). Synthesis, characterization, and enthalpy of mixing of the (Fe,Mg)CO<sub>3</sub> solid solution. *Geochem. Cosmochim. Acta.* 60, 4377–4383. doi:10.1016/s0016-7037(96)00261-x
- Daval, D., Martinez, I., Guigner, J.-M., Hellmann, R., Corvisier, J., Findling, N., et al. (2009). Mechanism of wollastonite carbonation deduced from micro- to nanometer length scale observations. *Am. Mineral.* 94, 1707–1726. doi:10.2138/ am.2009.3294
- Daval, D., Sissmann, O., Menguy, N., Saldi, G. D., Guyot, F., Martinez, I., et al. (2011). Influence of amorphous silica layer formation on the dissolution rate of olivine at 90°C and elevated pCO2. *Chem. Geol.* 284, 193–209. doi:10.1016/j. chemgeo.2011.02.021
- Effenberger, H., Mereiter, K., and Zemann, J. (1981). Crystal structure refinements of magnesite, calcite, rhodochrosite, siderite, smithonite, and dolomite, with discussion of some aspects of the stereochemistry of calcite type carbonates. *Z. Kristallogr.* 156 (1–4), 233–243. doi:10.1524/zkri.1981.156.14.233
- Fosbøl, P. L., Thomsen, K., and Stenby, E. H. (2010). Review and recommended thermodynamic properties of FeCO<sub>3</sub>. *Corrosion Eng. Sci. Technol.* 45, 115–135. doi:10.1179/174327808X286437
- Gadikota, G., Natali, C., Boschi, C., and Park, A.-H. A. (2014). Morphological changes during enhanced carbonation of asbestos containing material and its comparison to magnesium silicate minerals. J. Hazard Mater. 264, 42–52. doi:10.1016/j.jhazmat.2013.09.068
- Ghanbarian, B., Hunt, A. G., Ewing, R. P., and Sahimi, M. (2013). Tortuosity in porous media: a critical review. Soil Sci. Soc. Am. J. 77, 1461–1477. doi:10.2136/sssaj2012.0435
- Glynn, P. D., and Reardon, E. J. (1990). Solid-solution aqueous-solution equilibria; thermodynamic theory and representation. Am. J. Sci. 290, 164–201. doi:10. 2475/ajs.290.2.164
- Gunnarsson, I., and Arnórsson, S. (2000). Amorphous silica solubility and the thermodynamic properties of H4SiO<sup>°</sup>4 in the range of 0° to 350°C at Psat. *Geochem. Cosmochim. Acta.* 64, 2295–2307. doi:10.1016/s0016-7037(99)00426-3
- Hassal, P. (2008). Milling media review Part 2: bead density effect, Paint and coatings industry. Available at https://www.pcimag.com/articles/87925milling-media-review-part-2-bead-density-effect (Accessed June 27, 2020).
- Helgeson, H. C. (1969). Thermodynamics of hydrothermal systems at elevated temperatures and pressures. Am. J. Sci. 267, 729–804. doi:10.2475/ajs.267.7.729
- Huijgen, W. J. J., Comans, R. N. J., and Comans, R. N. J. (2005). Mineral CO<sub>2</sub> Sequestration by steel slag carbonation. *Environ. Sci. Technol.* 39 (24), 9676–9682. doi:10.1021/es050795f

Jankovic, A. (2000). Fine grinding in the Australian minerals industry. J. Min. Metall. 36 (1-2), 51-61.

- Julcour, C., Bourgeois, F., Bonfils, B., Benhamed, I., Guyot, F., Bodénan, F., et al. (2015). Development of an attrition-leaching hybrid process for direct aqueous mineral carbonation. *Chem. Eng. J.* 262, 716–726. doi:10.1016/j.cej.2014.10.031
- Knauss, K. G., and Wolery, T. J. (1988). The dissolution kinetics of quartz as a function of pH and time at 70°C. *Geochem. Cosmochim. Acta.* 52 (1), 43–53. doi:10.1016/0016-7037(88)90055-5
- Kwade, A., and Schwedes, J. (2007). Chapter 6: wet grinding in stirred media mills. *Handbook of Powder Technology*. 12, 251–382. doi:10.1016/s0167-3785(07) 12009-1
- Lee, S. H., and Rasaiah, J. C. (2011). Proton transfer and the mobilities of the  $H^+$  and  $OH^-$  ions from studies of a dissociating model for water. *J. Chem. Phys.* 135 (12), 124505. doi:10.1063/1.3632990
- Levenspiel, O. (1998). Chemical reaction engineering. 3rd Edn. New York, NY: John Wiley & Sons.
- National Academies of Sciences Engineering, and medicine (2019). "Chapter 3: mineral carbonation to produce construction materials," in *Gaseous carbon waste streams utilization: status and research needs*. Washington DC: The National Academies Press. doi:10.17226/25232
- Oelkers, E. H., and Schott, J. (2001). An experimental study of enstatite dissolution rates as a function of pH, temperature, and aqueous Mg and Si concentration, and the mechanism of pyroxene/pyroxenoid dissolution. *Geochem. Cosmochim. Acta.* 65 (8), 1219–1231. doi:10.1016/s0016-7037(00)00564-0
- Parkhurst, D. L., and Appelo, C. A. J. (1999). "User's guide to PHREEQC version 2—a computer program for speciation, batch-reaction, one-dimensional transport, and inverse geochemical calculations," in *Water-resources investigations report 99-4259*. Reston, VA: U.S. Geological Survey, 312. doi:10.3133/wri994259
- Parkhurst, D. L., and Appelo, C. A. J. (2013). Description of input and examples for PHREEQC version 3—a computer program for speciation, batch-reaction, onedimensional transport, and inverse geochemical calculations. Reston, VA: U.S. Department of the Interior, U.S. Geological Survey. doi:10.3133/tm6a43
- Parkhurst, D. L. (1995). User's guide to PHREEQC A computer program for speciation, reaction-path, advective-transport, and inverse geochemical calculations, Issues 95-4227. Reston, VA: U.S. Department of the Interior, U.S. Geological Survey, 143. doi:10.3133/WRI954227
- Pitzer, K. S. (1973). Thermodynamics of electrolytes. I. Theoretical basis and general equations. J. Phys. Chem. 77(2), 268–277. doi:10.1021/j100621a026
- Ploquin, F., Juillot, F., and Guyot, F. (2016). Caractérisation minéralogique et cristallochimique des produits de carbonatation des scories SLN et KNS. Programme « Carboscories », CNRT « Nickel & son environnement », internal report IMPMC. 50.

- Rimstidt, J. D. (1997). Quartz solubility at low temperatures. *Geochem. Cosmochim. Acta.* 61, 2553–2558. doi:10.1016/s0016-7037(97)00103-8
- Sanna, A., and Maroto-Valer, M. (2017). CO<sub>2</sub> sequestration by ex situ mineral carbonation. Singapore: World Scientific publishing. doi:10.1142/q0046
- Schilde, C., Kampen, I., and Kwade, A. (2010). Dispersion kinetics of nano-sized particles for different dispersing machines. *Chem. Eng. Sci.* 65, 3518–3527. doi:10. 1016/j.ces.2010.02.043
- Schott, J., and Berner, R. A. (1983). X-ray photoelectron studies of the mechanism of iron silicate dissolution during weathering. *Geochem. Cosmochim. Acta.* 47, 2233–2240. doi:10.1016/0016-7037(83)90046-7
- Sipilä, J., Teir, S., and Zevenhoven, R. (2008). Carbon dioxide sequestration by mineral carbonation. Literature review update 2005-2007. Report from Åbo Akademi University, Faculty of Technology, Heat Engineering Laboratory. Available at: http://innovationconcepts.nl/res/literatuurGPV/mineralcarbonation literaturereview0507.pdf (Accessed November 3, 2020).
- Sissmann, O., Brunet, F., Martinez, I., Guyot, F., Verlaguet, A., Pinquier, Y., et al. (2014). Enhanced olivine carbonation within a basalt as compared to singlephase experiments: reevaluating the potential of CO<sub>2</sub> mineral sequestration. *Environ. Sci. Technol.* 48 (10), 5512–5519. doi:10.1021/es405508a
- Touzé, S., Augé, T., Bodénan, F., Benhamed, I., Julcour, C., Bourgeois, F., et al. (2015). Carbonatation minérale en Nouvelle-Calédonie. Rapport de synthèse, Programme « Carboscories », CNRT Nickel & son environnement.
- Van Essendelft, D. T., and Schobert, H. H. (2009). Kinetics of the acid digestion of serpentine with concurrent grinding. 1. Initial investigations. *Ind. Eng. Chem. Res.* 48 (5), 2556–2565. doi:10.1021/ie801085x
- Veetil, S. P., and Hitch, M. (2020). Recent developments and challenges of aqueous mineral carbonation: a review. Int. J. Environ. Sci. Technol. 17, 4359–4380. doi:10.1007/s13762-020-02776-z
- Zhu, C., and Anderson, G. (2002). Environmental applications of geochemical modeling. Cambridge, UK: Cambridge University Press.

**Conflict of Interest:** 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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