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Efficiency of high energy over conventional milling of granulated blast furnace slag powder to improve mechanical performance of slag cement paste

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This work aims at bridging the efficiency of ball milling of granulated blast furnace slag (GBFS) to the structural and mechanical properties of slag cement pastes. Both conventional and high energy milling of GBFS are considered with a milling duration varied between 1 and 10 h. X-ray diffraction, infra-red spectroscopy, granulometry analysis and scanning electron microscopy are used to draw the main lines of structural and morphological changes occurring during milling. Cement pastes formulated using 45% of GBFS in substitution are characterized. Workability, X-ray diffraction analysis, differential scanning calorimetry and compressive testing are performed to analyse main structural changes and reactions driven by the presence of milled GBFS as well as its direct consequence on the mechanical strength of slag cement pastes. Slag milling indicates the superior efficiency of high energy milling, which allows a maximum slag fineness of 1.79 m²/g after 3 h of milling. Major structural changes occur during the first 3 h of high energy milling while conventional milling does not induce any remarkable trend. These changes concern amorphisation of the bulk structure in addition to the fracturing and agglomeration of slag particles. Workability of slag cement pastes is remarkably improved when using 1 h of high-energy slag milling. This result is consistent with slag fineness trend with respect to milling time and with the improvement of GBFS reactivity. The substitution of 45% of cement (CEM I 52.5) by GBFS is only beneficial at the condition of performing high-energy milling for at least 1 h.

Keywords:
Granulated blast furnace slag
High-energy ball milling
Cement paste
Compressive strength
Structural characterisation

1. Introduction

In extractive metallurgy, blast furnace slag (BFS) forms as a result of melting of iron oxides (raw ore, pellet, sinter) in blast furnace. Besides its role during iron smelting for assisting impurity removal from the ore, slag is used as an industrial by-product in several applications. In civil engineering, slag invaded two major markets, namely cement production and road construction. In particular, the market uptake justified major research direction for slag as a cement additive. For instance, Bellman and Stark [1] report the beneficial effect of slag on mechanical performance of slag mortar at an early age subject to appropriate activation. In chapter one of a book dedicated to by-products, Siddique [2] promotes the benefits of using Ground Granulated Blast Furnace Slag (GGBFS) in cement. More than a dozen of good reasons are exposed there such as good workability, improved strength and durability, resistance to sulphate attack, etc. Slag beneficial role on environment is reported by different contributors [3–5]. The use of slag as substitute is expected to reduce the environmental footprint of clinker manufacturing with a rate as large as five times less of CO₂ emission. This kind of substitution contributes to sustainable cement production since the construction sector is recognized as a large source of carbon emissions [6].

Different recycling routes of slag appear in the literature like the hydrothermal hot pressing explored by Yanagisawa et al. [7]. Their study shows that both pressure and water content are major influential process variables on tensile strength and bulk density of slag compacts compared to the reaction time.

Ozbay et al. [3] expose some critical factors, which act as leverage of slag hydraulic activity and heat of hydration. These are related to the chemistry of the slag, its degree of fineness and substitution ratio. The chemistry of the slag can be adjusted using activators such as NaOH.
Activators play a central role in producing CSH phases. Their concentrations modulate the degree of reactivity of slag as shown by Hilbig and Buchwald [8]. Complex hydration mechanisms take place during slag activation. These are responsible for the cleavage of the slag network and production of several mineral phases [9]. The understanding of the hydration mechanisms is important to anticipate reactivity of slag with cementitious materials. Soft X-ray transmission microscopy shows that development of hydration products in terms of morphology and growth rate is affected by the nature of activation suspensions [10].

According to the BFS cooling process, several types of slag can be structurally distinguished. Slag may appear as amorphous, crystalline or some combination of both [7,10–14]. If cooling is slow such as in free air, slag is grey, porous and exhibits a crystallized structure. However, if cooling is fast, such as when using water quenching [11], slag becomes vitrified.

Slag is referred as granulated slag when subject to grinding according to the standard NF IN 15167-1 [15]. Granulated blast furnace slag (GBFS) has a morphological form of clear-colour sand. GBFS shares the same main chemical compounds (mainly SiO₂, CaO, Al₂O₃, MgO) [2,11,16–20] as ordinary Portland cement [20–23]. GBFS has a direct utilization as additive in cement with rates varying from 6% up to 95% for CEM II and CEM III according to the norm NF EN 197-1 [22,23].

Mechanical strength improvement of slag concrete is not subject to major debate but a large discrepancy is observed for reported gains [11,24]. Jelidi et al. [11] show that only minor gain in compressive strength (~4%) is expected for slag concrete after 28 days. Concrete mixtures with GGBS allow only 15% of gain in long term (one year) compressive strength according to the work of Selem et al. [24].

The gain in mechanical performance is a challenging issue since the compressive strength of cement containing ~ 95% of clinker (CEM I) is nearly 53 MPa after 28 curing days [22,25]. Compared to other cement types containing high rates of slag such as CEM II (6%–35%) or CEM III (36%–95%), compressive strength decreases by 19% to 38%.

The challenge of increasing mechanical performance of slag cementitious materials is addressed in different ways. El-didamony et al. [20] considered different combinations of Portland cement, GBFS (ground granulated blast furnace slag) and drinking water sludge in the formulation of cement pastes. Compressive strength as large as 90 MPa is achieved for those pastes containing higher contents (~24%) of GBFS after 30 days of curing. The strategy of using additional phases to further increase the strength does not often pay off. Selem et al. [24] show that multi-phase mixtures involving GGBS (ground granulated blast furnace slag) is not suitable to for long term strength degradation under seawater attack. Their results indicate that GGBS alone is the most efficient additive against strength deterioration with only 3% of strength decrease after a year of exposure to sea water environment. Bellman and Stark [1] show that overcoming low early stage compressive strength of mortar is possible if hydration of slag is better improved. The authors show that the use of different salts resulting in the increase of pH improves the compressive strength of slag mortar by ~166% after 2 days of curing. The gain in strength represents 47% at an age of 28 days. Abdalqader et al. [26] considered alkali activation of fly ash/slag to improve the performance of cement using sodium carbonate. The best compressive strength trend with curing time is achieved when 10% of Na₂CO₃ is added to GGBS with 80 MPa as the highest score after 90 days.

GGBS reactivity with water under alkali environment as well as with calcium hydroxide determines the strength trend at early age upon the formation of hydration products [2]. Modulation of this reactivity using activation is certainly a promising pathway for improving strength of slag cementitious materials. Numerous ways of activation are reported in the literature, most of them rely on alkali activation [8,9]. Salman et al. [27] studied the effect of Na-silicate, KOH and NaOH activation of steel slag mortar. They achieved substantial rate of improvement of early age compressive strength (after 3 days of steam curing) with Na-silicate-NaOH compared to K-silicate-KOH activation. Despite the statement of the authors on the relatively moderate gain in strength (maximum strength 43 MPa) at long term curing (90 days), the reading of their results still indicate beneficial effect of alkali activation.

Both calcium sulphate addition [1] and chemical activation are also privileged ways to use efficiently slag in cementitious materials [15]. Dimitrova et al. [13] studied slag activation towards metal ions using thermal treatment. They demonstrate that copper removal is positively correlated to the slag heating temperature. Grinding techniques are also used as mechanical activators [28–33]. These techniques increase slag fineness, its specific surface area and thus they are expected to significantly affect pozzolanic and hydration kinetics. Wan et al. [28] used different types of grinding including ball, airflow and vibromill to relate GGBS fineness to strength of slag mortar. Highest fineness scores (i.e., smallest average diameter and largest surface area) are attributed to vibromill processing. The authors show that this grinding process results in largest compressive and flexural strengths of slag mortars.

Kriskova et al. [29] studied the effect of bead milling on mechanical activation of steel slag. Besides the substantial increase in surface area and amorphisation of slag, their study show that slag mortars exhibit different curing trends depending on the type of slag. However, mechanical activation demonstrates insufficient gain in both compressive and flexural strengths. Kumar et al. [30] introduced surface activation criterion in addition to specific surface area as a factor influencing mechanical activation efficiency of slag. The authors show that surface activation potential is nonlinearly correlated to milling time for maximum duration of 60 min [30]. Significant modulation of the hydration kinetics is obtained from non-reacted to complete hydrated slag. The result of such activation on mechanical performance of cementitious materials is, unfortunately, not considered by the authors. Allahverdi and Mahinroosta [31] attempted such study in the case of high phosphorous slag cement. The authors achieve compressive strengths as large as 120 MPa for slag mortar at an age of 180 days. This substantial improvement of compressive strength can be related to the large milling durations attempted by the authors (up to 18 h). It appears also from the results of the authors that the intensity of the milling reflected by the fineness is monotonously correlated to mechanical strength of slag mortar. Bougara et al. [32] combined similar finesses (between 250 and 420 m²/kg) of granulated blast furnace slag with alkali activation (using NaOH and KOH). Compressive strength results achieved by the authors are below 50 MPa at an age of 90 days and for slag content up to 50%. The authors considered also the effect of curing temperature between 20 °C and 60 °C [32]. They noticed the reduction of compressive strength with the increase of curing temperature even if they recommend the use of 40 °C as a curing temperature for an optimal strength increase trend. Behim and Clastres [33] correlated El Hadjar slag content (0–100%) and fineness (2500–4000 cm²/g) to the compressive strength of slag mortars. The largest compressive strengths are related to the largest finesses and smallest content of slag (10%). Strength scores <56 MPa are achieved for the best formulations after 90 days of hydration.

With classical or conventional grinding slag fineness does not exceed 5000 cm²/g after 24 h of milling [32,33]. High energy ball milling allows superior results (up to 9700 cm²/g in [29] and 8000 m²/g in [30]).

The analysis of the literature work on mechanical activation of slag cementitious materials shows that sparse information is provided about the structural justification of mechanical strength trends of slag cementitious materials. With no intention to diminish the quality and significant outcome of the research work attempted by several contributors, the focus of some of them on morphological analysis is obvious. This is assuredly related to the direct consequence of grinding on particle size and specific surface area analysis [28,32]. Some other contributors addressed the result of slag grinding process through various instrumental techniques such as XRD, calorimetry and infra-red (FTIR) analysis but bounded their interest to slag rather than slag in cementitious materials [29]. And some others were able to correlate the result of low energy grinding to structural analysis (XRD, calorimetry, TEM, FTIR) [30,31]. In this work, we propose to address this gap in structural
analysis of high energy ball milling activation of granulated blast furnace slag as well as its consequence on mechanical performance of slag cement pastes. The mechanical activation is considered here through high energy planetary ball mill thanks to friction mode process described in an earlier work \[34\]. The outcome of high energy milling is compared to classical milling of GBFS to highlight main differences. Substitution of cement CEM I by GBFS with a ratio as large as 45% is attempted. Knowing that CEM I contains large content of clinker (up to 95%), this substitution is intended to formulate more sustainable cement. This work addresses the intensity of pozzolanic reactions, which are expected to be modulated by grading parameters of GBFS.

2. Materials and methods

The major raw materials are cement CEMI 52.5 CP1 NF EN 197-1 from Calcia and granulated blast furnace slag provided by ArcelorMittal Algeria, SPA. The chemical composition of milled granulated blast furnace slag (GBFS) is performed using X-ray fluorescence spectroscopy. These experiments are carried out using the S2 RANGER instrument from Bruker and results are shown in Table 1. The chemical composition is also monitored during slag milling. This milling is carried out using both high energy and classical ball milling machines. High energy milling is based on planetary high-energy ball mill RETSCH PM400 from Retsch Company whereas conventional milling is based on Eberhard Bauer apparatus operating at a speed of 50 rpm. The milling is performed using steel balls having a diameter between 20 mm and 50 mm in diameter for duration of 10 h. The capacity of the milling conventional equipment is 0.05 m³, which allows processing 2 kg of slag in a single shot. High energy milling apparatus is composed of four vials mounted on a planar disc. The rotation of the disc is adjusted to 400 rpm. Vials rotation is in the opposite direction with a speed of 800 rpm. The milling time is varied in the range (1–9) hours. Grinding energy is transmitted to the slag using steel balls of 30 mm in diameter. The steel vial capacity is 500 ml. The weight of powder samples is adjusted to 200 g per vial whereas the ball-to-powder weight ratio is 4. Sealing is operated to avoid air contamination during milling process. Metal contamination is also managed by imposing a delay of 30 min after each hour of milling. This contamination emerges from iron or chromium release from the balls, vials or both.

Formation of the reference cement paste is performed using mixture of cement (C) and water (W) with W/C content ratio of 0.3. Slag cement pastes are prepared with a substitution ratio of 45 wt.%. Mixture of cement (C), grinding slag (GS) and water (W) is prepared using a constant proportion throughout the formulations with a ratio W/L of 0.3 where L = C + GS.

Cement paste mixtures are poured into right-prism moulds of dimensions 4 × 4 × 16 cm³. Samples are unmould after 24 h, cut into 3 cubes of 4 × 4 × 4 cm³ and stored in water vats at temperature of 20 °C. X-ray investigation is performed using Bruker D2 Phaser diffractometer with a continuous scanning mode and Cu Kα radiation (λ = 0.1541 nm). The lines are measured in the 2θ range (5–100)° by an increment of 0.02° for 15 s. The software used for building the X-ray diffraction diagrams is DIFFRAC.EVA with ICD PDF2. Sampling from the same vial is performed to achieve reliable X-ray diffraction diagrams as function of milling time. Sampling is repeated four times for the slag powders and two times for cement pastes. The change in lattice parameters of calcium hydroxide (portlandite) Ca(OH)₂ is calculated from the fitting of X-ray patterns using WinXRF software. This fitting considers the shift of the high angle diffraction line for all X-ray patterns using Bragg’s law. The crystallite size D is calculated using the XRD profile analysis determined by Williamson–Hall method \[35\]. More details concerning calculation method are available in \[36\]. IR spectroscopy is performed at room temperature using Thermo Scientific Nicolet IS10 instrument equipped with Smart iTR (with Diamond Plate) accessory. The spectra are processed using Omnic software.

Granulometer Laser LS 230 is used to determine particle size distributions within the range (0.04 μm–200 μm), Brunauer–Emmett–Teller (BET) adsorption method and Gemini VII 2390 Micrometrics instrument are considered to measure the specific surface area of milled slag. Due to limits of the apparatus, unmilled slag is sieved to achieve particle size smaller than 300 μm. Analysis of specific surface area is performed using static volumetric technique working on the principle of gas pressure balance. The internal volume and the temperature surrounding both tubes are maintained at identical conditions. The morphology of unmilled and milled powders are characterized using scanning electron microscopy (SEM) EVO40 (Carl Zeiss®) with a Bruker energy-dispersive spectrometer (EDS). The enthalpy change ΔH is determined using Differential Scanning Calorimetry based on NETZSCH DSC 04 F1 Phoenix equipment. Samples of about 20–25 mg are sealed in aluminum crucibles and heated at a rate of 20 °C/min under a nitrogen flow rate of 20 ml/min in the sample chamber and 70 ml/min rate in the furnace chamber. The total heat is measured in a wide temperature range (25–600 °C).

Mechanical testing is carried out using 3R 250 kN equipment. Compression testing is performed on cement paste samples of dimensions 4 × 4 × 4 cm³ at different curing times (7, 28 and 120) days. For each milling condition, 3 replicates are considered.

Workability of fresh slag cement paste is investigated using flowable test according to the NF EN 1015-3 norm \[37\]. The flow test uses a standard conical frustum shape with a diameter of 10 cm. The mould is centered on the flow table and filled in a two stage process with fresh cement paste. It is lifted vertically and tempered 15 times during 15 s. The relative change in flow of the cement paste is monitored to achieve flow or consistency evolution ε (%) following the expression:

\[
ε(\%) = \frac{(d - d_0)}{d_0}
\]

where d is the diameter of the cement paste after 15 times tamping, d₀ = 10 cm is the initial diameter.

3. Results and discussion

3.1. Morphology and structural properties of milled slag

Table 1 shows the results of chemical composition monitoring of GBFS as function of milling duration using high-energy ball milling. Also are reported the composition of the unmilled and the milled GBFS using the conventional milling. Table 1 confirms no traces of iron or chromium contamination. The composition of the as-received slag

<table>
<thead>
<tr>
<th>t (hour)</th>
<th>CaO</th>
<th>SiO₂</th>
<th>MgO</th>
<th>Al₂O₃</th>
<th>MnO</th>
<th>SO₃</th>
<th>BaO</th>
<th>Fe₂O₃</th>
<th>K₂O</th>
<th>SrO</th>
<th>Cl</th>
<th>TiO₂</th>
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<tbody>
<tr>
<td>0</td>
<td>48.04</td>
<td>30.8</td>
<td>6.58</td>
<td>5.57</td>
<td>3.16</td>
<td>1.82</td>
<td>0.7</td>
<td>1.01</td>
<td>0.9</td>
<td>0.38</td>
<td>0.168</td>
<td>0.158</td>
</tr>
<tr>
<td>1</td>
<td>47.59</td>
<td>29.84</td>
<td>6.6</td>
<td>5.81</td>
<td>2.59</td>
<td>1.72</td>
<td>0.66</td>
<td>1.08</td>
<td>0.9</td>
<td>0.29</td>
<td>0.15</td>
<td>0.28</td>
</tr>
<tr>
<td>3</td>
<td>47.73</td>
<td>30.7</td>
<td>4.6</td>
<td>5.51</td>
<td>2.31</td>
<td>2.08</td>
<td>0.49</td>
<td>1.03</td>
<td>0.88</td>
<td>0.33</td>
<td>0.13</td>
<td>0.26</td>
</tr>
<tr>
<td>6</td>
<td>48.42</td>
<td>30.17</td>
<td>5.9</td>
<td>5.70</td>
<td>2.64</td>
<td>1.53</td>
<td>0.69</td>
<td>1.04</td>
<td>0.77</td>
<td>0.30</td>
<td>0.12</td>
<td>0.30</td>
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<tr>
<td>9</td>
<td>47.24</td>
<td>29.62</td>
<td>6.9</td>
<td>5.85</td>
<td>2.44</td>
<td>1.63</td>
<td>0.64</td>
<td>1.01</td>
<td>0.86</td>
<td>0.27</td>
<td>0.11</td>
<td>0.23</td>
</tr>
<tr>
<td>10⁵</td>
<td>48.15</td>
<td>30.14</td>
<td>6.52</td>
<td>5.91</td>
<td>3.01</td>
<td>1.92</td>
<td>1.05</td>
<td>1.09</td>
<td>0.91</td>
<td>0.24</td>
<td>0.14</td>
<td>0.20</td>
</tr>
</tbody>
</table>

* Conventional milling.
contains a typical high rate of lime (CaO), a large content of silica (SiO₂), a low content of alumina (Al₂O₃) and magnesia (MgO). These are the most representative species of the studied slag as they explain 94% of the composition. The unmilled slag is also typical to what is reported in the literature about blast furnace slag aggregate [2,11,16–20]. Only slight changes in the oxide contents are not to be mentioned.

The effect of milling on slag composition is rather minor. Major oxides composing the slag for which the content is larger than 5% do not exhibit significant change (almost 10% of variation) if we exclude magnesia (MgO) content which shifts from 6.58% down to 4.6% after 3 h of milling (30% of variation). Furthermore, manganese oxide (MnO) content change oscillates between 16% (content of 2.64% at t = 6 h) and 27% (content of 2.31% at t = 3 h) with no consistency of the trend with the milling time.

Fig. 1 compares XDR patterns of processed GBFS using both conventional and high energy ball milling. The spectra relative to high energy milling are differentiated using the milling time (t). XRD pattern corresponding to t = 0 h refers to the pattern of the as-received GBFS. This latter pattern shows diffuse scattering signature of the amorphous GBFS structure identified from the broad diffuse hump peak in the interval 2θ between 20° and 40°. This pattern is reported in the literature as a typical characteristic of the vitreous structure of GBFS due to the presence of glassy fractions [17,19,33,38]. The XRD pattern of the as-received GBFS shows the presence of vaterite mineral form of CaCO₃ identified from the peaks (112) and (008) at 2θ = 27.1° and 42.8°, respectively. XRD pattern of GBFS similar to the one studied here issued from (same provenance than our GBFS), Behim et al. [33] report XRD (CoKα radiation) spectra of El-Hajar GBFS similar to the one studied here. The authors identify the two diffuse hump peaks at approximately 2θ = 20°–48° and 2θ = 46°–68°. The same authors [33] report crystallized minerals like calcite, iron and traces of gehlenite (C₃AS) or akermanite (C₂MS₂) within the amorphous bulk. XRD patterns of GBFS from Taiyuan steel company in China [9] issued using CuKα irradiation places the broad diffuse hump peak in the region of 2θ between 20° and 38°. Zhang et al. [9] shows the presence of four kinds of mineral phases, namely akermanite, gehlenite, calcium silicate, and merwinite.

Structural modifications inferred to conventional milling can be read from Fig. 1. We observe a shift by 3° of the diffuse hump peak to the range 2θ (23°–35°). This shift is accompanied by the appearance of the mineral calcite containing the most thermodynamically stable form of CaCO₃. Calcite is identified by the peak (104) at 2θ = 29.4°. As for high-energy ball milling, complete amorphisation of the bulk material is achieved after the first hour of milling. Both conventional and high-energy milling have comparable diffuse scattering signature. The difference between the two milling processes comes from the large milling duration (10h) required by conventional milling to achieve the same amorphisation degree resulting from only 1 h of high energy ball milling.

The superior result of high-energy milling is explained by the intense nature of the repeated welding and fracturing of the powder mixture. During high energy milling, severe plastic deformation process of the powder particles takes place stimulated by ball-particle-ball and ball–particle vial wall interactions [39,40]. These results compare fairly with the general trend of amorphisation reported by Kriskova et al. [29] on slag bead milling for 6 h. The authors [29] observe also differences between minerals in terms of degree of amorphisation.

Fig. 2 compares infrared spectra of GBFS using both ways of milling. Assignment of peak bands of the as-received GBFS is provided by the

![Fig. 1. XRD patterns of GBFS using conventional and high energy millings.](image1)

![Fig. 2. FT-IR spectra of GBFS using conventional and high energy millings.](image2)
spectral search feature implemented in the Thermo Scientific Omnic series software. The search performed with a sensitivity of 80% delivers the assignments summarized in Table 2. These are also highlighted by dash lines in Fig. 2. All OH groups identified in the band width 4000–2500 cm\(^{-1}\) for the as-received slag show up for all milling conditions but with slight shifts and smaller intensities. In the range 2500–2000 cm\(^{-1}\), C=O and H–O–H stretching modes are present in the raw material. These also resist both conventional and high energy milling except at the position 2017 cm\(^{-1}\). This band peak disappears after both types of milling. In the range 2000–500 cm\(^{-1}\), the band position at 1419 cm\(^{-1}\) is assigned to the symmetric stretching mode of the O–C=O bond (Table 2). This stretching bond is identified at a different position (1450 cm\(^{-1}\)) by Abdalqader et al. [26]. Conventional milling shifts the peak form the band position 1419 cm\(^{-1}\) to 1437 cm\(^{-1}\) with a decreasing intensity. This shift can be attributed to the presence of different form of CaCO\(_3\) as discussed from the XRD patterns (Fig. 1). The same band peak disappears after the first hour of high energy ball milling due to the complete amorphisation of the bulk material.

Si–O stretching band is identified in the as-received GBFS by two peaks at 874 cm\(^{-1}\) and 1004 cm\(^{-1}\). These peaks are replaced by one peak at 911 cm\(^{-1}\) highlighted using a bold dash line after milling (both conventional and high-energy ball milling). Abdalqader et al. [26] attribute the shoulder at 875 cm\(^{-1}\) to the asymmetric stretching of Al\(_2\)O\(_3\) groups. A minor sharp peak at the position 713 cm\(^{-1}\) is assigned to Ca–O stretching band. This peak is shifted to smaller wavenumbers and flattens upon both millings. The last identified band peak at position 662 cm\(^{-1}\) corresponds to Si–O–Si bending vibration modes. Due to its relative weak intensity, this peak is not highlighted in Fig. 2. The associated bending mode resists milling for all attempted durations.

Particle size distribution analysis of GBFS is presented in Fig. 3 for all milling conditions. Fig. 3a exhibits the main modifications occurring on the particle size profiles depending on milling type and duration. All particle size distributions are plotted in cumulative form. Due to the large-size particles composing the as-received slag powder, granulometry is determined using classical sieving method. The median size value D\(_{50}\) of the as-received slag is close to 1.1 mm. Granulometry of 85% of the distribution is below a size of 2 mm. If laser diffraction is used to obtain the initial particle size of the as-received slag, this leads certainly to underestimation of the outcome of the milling process on particle size reduction. Salman et al. [27] used laser granulometry to characterize the size dispersion of stainless steel refining slag. They achieve a median value of 45 \(\mu\)m and point out the limit of the technique to appropriately address shape effects. The reading of their SEM micrograph shows also possible underestimation of the starting granulometry.

In the present study, a size shift of about two orders of magnitude appears as a result of milling (Fig. 3a). Granulometry achieved using conventional milling is close to the one obtained at larger high energy milling durations (\(t > 3\) h). However, the kinetics of particle size shrinkage related to high energy milling shows that milling types are far from being equivalent. Indeed, significant particle size reduction is achieved after the completion of the first hour of high energy milling. The further increase in particle size for larger times (\(t > 1\) h) demonstrates an evidence of particle agglomeration. This kinetics is further described later using SEM images. Quantitative analysis of size class trends as function of milling time shows the following (Fig. 3b). Evidence of agglomeration is supported by the D15 size class. Fine particles (D15) shows a steep increase in size up to 6 h of high energy milling with an average size of 2.5 \(\mu\)m reached at \(t = 6\) h. The D15 trend surpasses the achieved particle size using conventional milling. However, the analysis of the final milling duration shows that there is some transfer to the upper class (D60), which results in a smaller particle size. The same trend is noticed for D60 but with a smaller rate. Here, the transfer to upper classes due to agglomeration is limited compared to the previous case. The largest D60 value remains smaller compared to the average size achieved by conventional milling. The same trend is noticed for D85 with a slower rate of particle size increase with respect to milling duration. Finally, the D100 trend is characterized by a strong competition between particle fracturing and agglomeration. In particular, particle shrinkage continues up to an optimal milling time of 3 h. The monotonous positive trend noticed for \(t > 3\) h translates the upper hand of agglomeration over particle shrinkage.

In order to keep going on fracturing of small particles (<30 \(\mu\)m), 1 h or less of high energy milling is optimal. Fracturing of larger particles can be considered up to 3 h but a cost of accepting agglomeration of smaller ones. Kumar et al. [30] considered attrition milling of slag up to 1 h. The authors show a continuous shift of particle size distribution

![Fig. 3. (a) Particle size cumulative distribution as function of milling conditions. (b) Analysis of average particle size versus milling time for different particle size classes.](image)

Table 2
Assignments of FT-IR peaks corresponding to the as-received slag.

<table>
<thead>
<tr>
<th>Wave number (cm(^{-1}))</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>662</td>
<td>Si–O–Si bending vibration modes</td>
</tr>
<tr>
<td>713</td>
<td>Ca–O stretching band</td>
</tr>
<tr>
<td>874, 1004</td>
<td>Si–O stretching band</td>
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<td>1419</td>
<td>O–C=O asymmetric stretching vibration bonds</td>
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<tr>
<td>2017, 2160</td>
<td>H–O–H stretching</td>
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<td>2312</td>
<td>C–O stretching vibration</td>
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<tr>
<td>3667, 3814, 3851</td>
<td>OH groups</td>
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</table>
of slag towards small classes with a median value reaching 3.7 μm at the final milling time.

Results of slag milling reported by Kriskova et al. [29] show no evidence of agglomeration for large milling durations (between 2 and 6 h). With regards to the achieved specific surface area in their results, bead milling used in their study can be considered as intensive. This result contrasts with the present study. The main difference between their results and ours may be explained by the presence of process control agents (ethanol), which is supposed to enhance fracturing phenomenon.

Fig. 4 shows the evolution of GBFS morphology as function of milling duration. The as-received slag exhibits an airy irregular shape with a smooth surface. Although, it is difficult to measure characteristic dimensions from SEM, the surface density of porosities seems to be close to 4% (Fig. 4a). The porosity size varies between 3 and 88 μm with an average size of 10 μm. Result of conventional milling (Fig. 4b) shows significant particle size dispersion. In addition, particle morphology modification is evident upon milling. Larger particles maintain, to some extent, the irregular shape, whereas smaller ones appear more globular and sticky. The shape factor (i.e., ratio between the smallest and largest intercept length) is 0.51 ± 0.07 for large particles. The same structural attribute measured for small particles is close to 0.91 ± 0.17. No evidence of porosity is found within both small and large particles. The observed change in morphology reflects the result of repeated welding, fracturing, and rewelding of particles [40]. High-energy milling performed for 1 h (Fig. 4c) leads to the same particle size reduction observed for conventional milling but differences in particle morphology and degree of fines are striking. We observe a denser and homogeneous population of fine particles that decorate larger ones. The smaller particles tend to weld together and the larger particles to fracture. Further increase of high energy milling (Fig. 4c–f) accentuates particle rewelding to such a state where the milled powder appears as a connected cohesive network. The development of the particle network can be explained by an increase of the local temperature, which enhances cohesive forces between aggregates [41].

The balance between fracturing and welding seems to be achieved within the first hour of milling (Fig. 4c) as no evident of reduction in size of large particle is perceived for larger milling times. Wan et al. [28] confirm morphological changes from irregular to spherical shapes through SEM observations of milled slag. The authors claim also the possibility to control better the particle size dispersity using airflow mill and show different outcomes depending on the type of milling process.

To better capture the outcome of the present milling process, specific surface area results are exposed in Fig. 5. The specific surface area of the as-received slag is 0.29 m²/g. Most reported results in the literature places specific surface area levels for GBFS between 0.37 m²/g and 0.48 m²/g [2]. Conventional milling increases specific surface area by 38%. This is a much lesser extent than the result of the first hour of high energy milling (434%). In addition, the increase of specific surface area reaches its maximum value (1.8 m²/g) after 3 h of high-energy milling.

**Fig. 4.** SEM micrographs showing the evolution of slag particle morphology from the (a) as-received to the (b) conventional and (c) high energy milling for milling time (t) of (c) 1 h, (d) 3 h, (e) 6 h and (f) 9 h.
milling. The decreasing branch observed for longer milling time confirms the tendency of agglomeration. This result is supported by the analysis of granulometry (Fig. 3), and more particularly the trend exhibited by the D100 size class (Fig. 3b).

Reported results on specific surface area evolution as function of milling are numerous. Wan et al.[28] have found a lower range of specific surface area values between 0.51 m²/g and 0.685 m²/g for different milling processes. Kriskova et al.[29] have found higher levels of specific surface area in the range (0.7 m²/g–9.7 m²/g) after 6 h of milling.

3.2. Workability of cement-slag pastes

Slag cement pastes prepared using 45% of slag in substitution result in the workability tendencies shown in Fig. 6. One can read the similarity between consistency scores and specific surface area results shown in Fig. 5. The reference cement CEM I paste exhibits the lowest consistency score (ε). The use of conventional milling has an unperceivable effect on cement paste consistency. One hour of high energy is sufficient to increase the consistency of the slag cement paste by 43% to its maximum level. A more stable steady-state consistency is observed for longer milling times. These observations match the workability tendencies reported on cementitious materials modified using slag [2,3,28,42]. For instance, the same positive correlation between workability and specific surface area is pointed out by Wan et al.[32]. In addition, Nath and Sarker. [42] noticed that the slump of concrete and flow of mortar decreased with the increase of slag content in the mixtures.

3.3. Structural and thermal behavior of slag cement paste

X-ray diffraction patterns of slag cement pastes performed are shown in Fig. 7 for all milling conditions. The reference paste cement CEM I contains calcium hydroxide (portlandite) Ca(OH)₂ of a hexagonal structure characterized by different plan peaks P(001), P(100), P(101), P(102), P(110), P(111), P(201), P(112). We have also evidence of calcium carbonate (calcite) CaCO₃ presence with rhombohedral structure characterized by different plan peaks C(104), C(113), C(202). In addition, calcium silicate hydrate (CSH) is detected with Ca₂SiO₄·3H₂O monoclinic structure identified by S(100), S(112), S(016), S(-321), S(-221) and also ettringite with Ca₆Al₂(SO₄)₃(OH)₁₂·6H₂O hexagonal structure identified by E(101), E(220), E(304), E(322) and E(332). Former studies indicate that CSH gel is generally identified as an amorphous structure [43,44] and Ettreinigite is detectible after long curing time [45]. Paste cement CEM I modified by 45% of slag milled using conventional route exhibits the same collection of peaks of nearly the same intensities. In addition, the presence of peak S₁(211) is attributed to a second C-S-H with Ca₂SiO₄·H₂O orthorhombic structure. Other characteristic peaks A(102), A(522), A(260) are informative of the presence of C-A-H with Ca₂(AlO₂)₃(OH)(H₂O) orthorhombic structure.

Modified cement pastes with high energy milled slag shows evidence of C-S-H peak S₁(211) irrespective of milling duration (Fig. 7). The remarkable effect of high energy milling can be attributed to the decreasing intensity, enlargement and slight shift of portlandite peaks. The decaying portlandite peaks can be attributed to pozzolanic reactions between activated slag and calcium hydroxide. The hydration
process of Portland cement can be summarized as follows [46,47]:
\[
\begin{align*}
  C_2S + 2H &\rightarrow 0.5C_3S_2H_6 + 0.5CH \quad (1) \\
  C_2S + 3H &\rightarrow 0.5C_3S_2H_6 + 1.5CH \quad (2)
\end{align*}
\]

whereas the pozzolanic reaction can be summarized as flows [48,49]:
\[
\begin{align*}
  S + 1.5CH &\rightarrow 0.5C_3S_2H_6 \quad (3) \\
  A + 4CH + 9H &\rightarrow CaAH_{13} \quad (4)
\end{align*}
\]

where $C$: CaO, $S$: SiO$_2$, $A$: Al$_2$O$_3$, $H$: H$_2$O, $CH$: Ca(OH)$_2$.

As observed from the expressions (3) and (4), the active silica (SiO$_2$) and aluminate (Al$_2$O$_3$) of milled slag react with calcium hydroxide originated from Portland cement hydration giving raise to calcium silicate hydrate, i.e., $C$-$S$-$H$ like in expression (3) or calcium aluminate hydrate, i.e., $C$-$A$-$H$ like in expression (4). Shi and Day [50] have studied the products of pozzolanic reaction mechanisms in lime + pozzolan cement pastes in the presence of Na$_2$SO$_4$ and CaCl$_2$. Their X-ray diffraction (XRD) analysis pointed out the large and sharp diffuse band of calcium silicate hydrate ($C$-$S$-$H$) with curing time and the vanishing trend of Ca(OH)$_2$ after 180 days. We believe that, in our case, the boarding and slight shift of portlandite peaks can be attributed to the reduction of crystallite size and inter-reticular distance of portlandite, respectively [36].

Fig. 8 shows the evolution of lattice parameter ($a$, $c$), cell volume ($V = a \times a \times c$) and crystallite size of portlandite for all studied formulations. The lattice parameters ($a$) and ($c$) of portlandite structure given by ICDD file are $a = 0.3590$ nm and $c = 0.4916$ nm, respectively. The same parameters determined from the reading of XRD results show slight changes with $a = 0.3604 \pm 0.0005$ nm and $c = 0.4942 \pm 0.0005$ nm. The lattice parameter ($a$) decreases with the increase of high energy milling time up to $3$ h before stabilization. This result contrasts with the effect of conventional milling, where negligible variation of lattice parameter ($a$) is observed. The lattice parameter ($c$) follows the opposite trend versus milling time with the same stabilization after $3$ h of high energy milling. Here also, the effect of conventional milling is minor. The evolution of the cell ($V$) and the crystallite size ($D$) confirm the critical effect of the three first hours of high energy milling, more particularly for the crystallite size, $D$. Indeed, the crystallite size decreases from its reference condition ($D = 79.4 \pm 2.5$ nm) to its steady-state value ($D = 17 \pm 1.5$ nm), which represents about 79% as an amount of variation. In contrast to this, conventional milling has no particular effect on both crystallite size and cell volume.

Fig. 9 shows DSC results of slag cement pastes for all formulations involving slag milling time. Heat flow curves show the existence of some endothermic peaks attributed to the decomposition of water bonding in two particular regions. The first region of decomposition is bounded between $100^\circ$C and $350^\circ$C. It is a characteristic of adsorbed water and may be related to decomposition of ettringite, dehydration of calcium-silicate hydrate ($C$-$S$-$H$) gel and calcium-aluminate hydrate. This statement is based on identified phases from the XRD analysis (Fig. 8). The second stage of decomposition is in the range of $400^\circ$C and $550^\circ$C. This region is characteristic of crystallized water and corresponds to the decomposition of Ca(OH)$_2$. The confrontation of these observations with the concerned literature show that the range between $50^\circ$C and $300^\circ$C is generally attributed to decomposition of ettringite, $C$-$S$-$H$, $C$-$A$-$H$ and monosulfoaluminate, whereas the range from $400^\circ$C to $500^\circ$C is attributed to decomposition of calcium hydroxide [51–55].

To follow the enthalpy change ($\Delta H$) associated with the decomposition of both adsorbed and crystallized water, an estimate on the areas under the endothermic peaks in the temperature range ($100^\circ$C–$250^\circ$C) and ($400^\circ$C to $550^\circ$C) is provided. Table 3 summarizes ($\Delta H$) values as function of milling time. For cement paste formulations involving high energy milling ($0 < t < 10$ h), we observe that $\Delta H$ associated to the decomposition of crystallized water follows a two-stage evolution with a first increasing stage up to $3$ h and then a steady-state value. The decreasing tendency of $\Delta H$ associated to crystallized water can be related to the pozzolanic reactions between milled slag and calcium hydroxide. This explanation is supported by DRX results. The former trend is not ultimately observed for $\Delta H$ associated to absorbed water. In this case, a slight increase after $3$ h of high energy milling can be attributed to secondary $C$-$S$-$H$ formation from pozzolanic reactions between milled slag and Ca(OH)$_2$. Undetectable changes in both $\Delta H$ values associated to crystallized and absorbed waters for those cement paste formulations involving slag milling using conventional route.

![Fig. 8. Evolution of the lattice parameters ($a$, $c$), cell volume ($V$) and crystallite size ($D$) of portlandite as function of milling time for both conventional and high energy milling.](image)

![Fig. 9. DSC curves of slag cement pastes as function of milling time.](image)

**Table 3**

<table>
<thead>
<tr>
<th>Milling time, $t$ (h)</th>
<th>$\Delta H$ adsorbed water ($\mu$J/g)</th>
<th>$\Delta H$ crystallized water ($\mu$J/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>80.5 ± 2</td>
<td>115.2 ± 3</td>
</tr>
<tr>
<td>10$^a$</td>
<td>79.4 ± 1.7</td>
<td>104.5 ± 3</td>
</tr>
<tr>
<td>1</td>
<td>76.1 ± 2.1</td>
<td>68.7 ± 2.8</td>
</tr>
<tr>
<td>3</td>
<td>87.1 ± 2.4</td>
<td>55.7 ± 2.5</td>
</tr>
<tr>
<td>6</td>
<td>88.1 ± 1.8</td>
<td>54.8 ± 2.7</td>
</tr>
<tr>
<td>9</td>
<td>82.1 ± 1.7</td>
<td>55.2 ± 2.8</td>
</tr>
</tbody>
</table>

$^a$ Conventional milling.
Compression strength of slag cement pastes as function of milling and curing conditions. The best situation corresponds to 3 h of milling, which results in the largest gain (8.8%) in compressive strength for all curing days when conventional milling of slag is used in the formulation of cement pastes. The optimal situation corresponds to a strength loss of −14% after 120 of curing days. An overall evaluation of the effect of high energy milling of slag shows a beneficial role on compressive strength of cement pastes. The optimal situation corresponds to 3 h of milling, which results in the largest gain (8.8%) in strength after 120 curing days. This represents a compressive strength of 62 MPa.

4. Conclusions
High energy milling proves to be an effective way to achieve high rates of slag fineness. High energy milling efficiency is limited by the agglomeration of slag particles, which is observed for milling durations superior to 3 h. This study concludes that the most relevant milling durations are within the hour unit. The decrease of the crystallite size of portlandite seems to be related to the efficiency of high energy milling within the first 3 h of the process. In contrast, such behavior is not observed for conventional milling. Mechanical activation of slag in cement paste is demonstrated through the presence of C–S–H and the decaying profile of portlandite. High energy milling has a direct effect on the pozzolanic reactions that take place in the slag cement paste.

Mechanical results provide an insight on the substitution of cement by milled slag. Improvement of both short and long term compressive strength of slag cement paste is possible with formulations involving 45% of cement CEM I substitution with high energy milled slag.

Acknowledgments and funding information
We thank Margarita Walferrdein from IRC/ESTP, Cachan, France, for her technical assistance concerning structure and mechanical measurements.

References

Fig. 10. Compression strength gain of slag cement pastes as function of milling time.

Table 4
Compression strength of slag cement pastes as function of milling and curing conditions.

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Slag milling time (h)</th>
<th>Curing strength (MPa)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Curing, 7 days</td>
</tr>
<tr>
<td>CEMI000h</td>
<td>52 55 57</td>
<td>40 46 49</td>
</tr>
<tr>
<td>PCEM10H</td>
<td>47 54 60</td>
<td>52 57 63</td>
</tr>
<tr>
<td>PCEM5H</td>
<td>53 55 56</td>
<td>53 55 63</td>
</tr>
</tbody>
</table>

* Conventional milling.

Table 4
Compression strength of slag cement pastes as function of milling and curing conditions.

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