The Incorporation of Steel Slag into Belite-Sulfoaluminate Cement Clinkers

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Abstract: The potential use of steel slag from treated steel slag in belite-sulfoaluminate cements was investigated in this study. Cement clinkers with two phase compositions were synthesized, allowing the incorporation of different amounts of steel slag. The phase composition and microstructure of cement clinkers at three different sintering temperatures were studied by X-ray powder diffraction and the Rietveld method, as well as scanning electron microscopy with energy dispersive spectrometry. The results showed that the targeted phase composition of clinkers was achieved at a sintering temperature of 1250 °C. However, a higher amount of perovskite instead of ferrite was detected in the clinker with a higher content of Ti-bearing bauxite. Apart from the main phases, such as belite, calcium sulfoaluminate, and ferrite, several minor phases were identified, including mayenite, perovskite, periclase, and alkali sulfates. In both clinker mixtures, a higher content of MgO in the steel slags resulted in the formation of periclase. Furthermore, the hydration kinetics and compressive strength at 7 and 28 days were studied in two cements prepared from clinkers sintered at 1250 °C. As evidenced by the results of isothermal calorimetry, the hydration kinetics were also influenced by the minor clinker phases. Cement with a higher content of calcium sulfoaluminate phase developed a higher compressive strength.

Keywords: cements; BCSA; steel slag; secondary raw materials; microstructure; clinker phase

1. Introduction

The co-generated ferrous slag stream is an integral part of the steel production process. Slag is the biggest by-product stream of the steel sector, accounting for 90% with regards to the mass of by-product and waste generated [1]. In total, 30% of the world’s steel is produced using an Electric Arc Furnace (EAF), with scrap steel as the input and electricity as the energy source. On average, around 200 kg of co-products are generated from one ton of steel produced in this way [2]. In Europe, 15.7 million tons of steelmaking slag were produced in 2018, with 34.8% coming from EAF-based production and 12.5% from other processes, probably secondary metallurgical processes [3]. The most common use of EAF slag is as aggregate in road construction, for which EAF slag from carbon steel production is predominately used (EAF C slag). EAF C slag has excellent mechanical properties, which justify its use as an aggregate in the manufacturing of bituminous mixes [4]. Other uses for steel slags (EAF C slag, EAF stainless steel slag (EAF S slag), and ladle slag coming from secondary metallurgical processes) are as aggregate for concrete, waste water treatment, embankments and fill, and ground stabilization [5]. Their use in cement and other hydraulic binders has also been reported [6–9]. Although steel slag is highly recyclable, there is still room for improvement, and efforts in the steel industry are being made to move closer to the “zero-waste” goal. In order to achieve the “zero waste” goal, it is important to look for alternative applications for the use of steelmaking slags that are not yet in widespread use. In the cement industry, the possibility of producing low-carbon and low-energy cements using secondary mineral raw materials rich in aluminium
is a promising solution, which would save natural resources and reduce the environmental impact of the production processes. To date, only a few studies have demonstrated the use of steel slags in the production of belite-sulfoaluminate (BCSA) clinker [10,11]. BCSA clinkers are one of the most promising low carbon and low energy potential alternatives to ordinary Portland cement (OPC) clinkers, as they use a sintering temperature that is around 200 °C lower and less energy is consumed for the grinding process [12–14]. They consist of three main phases: Belite (C$_2$S); calcium sulfoaluminate (C$_4$A$_3$S); and ferrite (C$_4$AF). By definition, the most abundant phase is belite, which represents more than 50 wt.% of BCSA clinker [14]. Among the minor phases, periclase (M), gehlenite (C$_2$AS), mayenite (C$_{12}$A$_7$), perovskite (CT), arcanite (KS), acermanite, magnetite, and anhydrite (CS) are the most commonly reported [15,16]. Although the targeted amounts of the main clinker phases can be well-predicted, the formation of minor phases strongly depends on the minor elements which are present in the raw materials [17,18]. Moreover, foreign elements can be incorporated into the main clinker phases, influencing the dynamics of crystal formation [19–21] and consequently affecting the hydraulic properties of the cement [22–24].

The aim of this study is to demonstrate and discuss the use of steel slag in BCSA cement clinker. BCSA clinkers of two targeted phase compositions were sintered at 1200, 1250, and 1300 °C, in order to describe the general microstructure, phase composition, and incorporation of minor elements into the hydraulic clinker phases. To reveal any potentially unfavorable influence of the material-derived impurities on the hydraulic properties, BCSA cements were prepared from clinkers sintered at 1250 °C. Early age hydraulic properties were investigated by isothermal calorimetry and compressive strength development was measured at 7 and 28 days of hydration.

2. Materials and Methods
2.1. Materials

The main raw materials used in this study were limestone and flysch (deep water siliciclastic rocks), steel slag, calcined bauxite, and white titanogypsum. For the correction, smaller quantities of mill scale, quartz, and calcite were also used. Steel slag, or more precisely slag mineral residue, was obtained from the processing of the mixture of EAF S slag and ladle slag. Slag processing involved weathering in the slag cooling yard, grinding, and metal recovery. The final product of the slag processing was steel granulate, which was returned to the EAF as steel scrap and slag mineral residue, the latter of which was used in this study. The treatments and processes which gave the mixture of EAF S slag and ladle slag its final properties are normal industrial practices. These processes and treatments are reported as Best Available Techniques by BREF documents.

With the exception of white titanogypsum, which was dried at 40 °C to prevent its transformation to anhydrite, all raw materials were dried to a constant mass at 105 °C. In order to prepare large batches of homogeneous raw materials, 5 kg of each material was milled in a ball mill, until it met the criteria that 95 wt.% of the material could pass through a 200 µm sieve. The exception was gypsum, which was not milled. The main oxide composition and loss on ignition at 950 °C in the raw materials for the clinker are given in Table 1.

The proportions of raw materials in the raw clinker mixtures were calculated using the Bogue method for sulfoaluminate cements [25]. The formulation of targeted phase compositions was based on the minimum and maximum amount of C$_4$A$_3$S in belite-rich sulfoaluminate clinkers [26]. A medium amount of C$_4$AF was pointed (constant for both mixtures) and the C$_2$S content was adjusted to reach the sum 95 wt.% (minor phases were estimated to be 5 wt.%). The intended phase compositions are given in Table 2, and the corresponding proportions of raw materials in the raw mixtures KMO1 and KMO2 are presented in Table 3.

Raw mixtures were homogenized in a laboratory ball mill (CAPCO Test Equipment Ball Mill Model 9VS). In total, 200 g of each mixture was homogenized in 200 mL of
isopropanol, together with steel balls, for 3 h. Isopropanol was removed by vacuum drying (1 medium permeable filter paper was used), followed by oven drying at 40 °C overnight. The raw material mixtures were then pressed into pellets 30 mm in diameter, using a force of 10.6 kN, with a weight of 15 g. The clinker mixtures were sintered at 1200, 1250, and 1300 °C in a Protherm furnace (PLF 160/9), at a heating rate of 10 °C/min. Samples were held at the final temperature for 60 min and allowed to cool naturally in the closed furnace.

**Table 1.** Chemical composition of the raw materials (wt.%). The main oxides needed for the formulation of the raw mixtures were determined by wet chemistry, while the minor components were determined by X-ray fluorescence spectrometry (WDXRF, Thermo Scientific ARL PERFORM'X, Thermo Fisher Scientific, MA, USA; fused beads, UniQuant program).

|          | Limestone | Gypsum | Calcined Bauxite | Flysch | Steel Slag | Quartz | Calcite | Mill Scale |
|----------|-----------|--------|------------------|--------|------------|--------|---------|------------|
| LOI 950 °C | 41.41     | 21.35  | 0.15             | 24.65  | 5.02       | 10.00  | 42.80   | 0.00       |
| SiO₂      | 4.76      | 0.13   | 5.88             | 32.02  | 17.46      | 85.50  | n.a.    | 1.08       |
| Al₂O₃     | 0.86      | 1.16   | 87.69            | 7.74   | 9.18       | n.a.   | 95.75   | n.a.       |
| Fe₂O₃     | 0.51      | 0.04   | 1.89             | 3.49   | 10.58      | n.a.   | 0.20    | 0.01       |
| CaO       | 51.51     | 32.62  | 0.06             | 28.83  | 39.59      | n.a.   | 56.06   | 0.19       |
| MgO       | 0.90      | 0.04   | 0.52             | 1.69   | 12.16      | n.a.   | n.a.    | 0.29       |
| SO₃       | 0.09      | 45.39  | 0.17             | 0.07   | 0.36       | n.a.   | 0.30    | 0.01       |
| Na₂O      | 0.81      | 0.26   | 0.08             | 0.54   | 0.21       | n.a.   | 0.67    | n.a.       |
| K₂O       | 0.14      | 0.01   | 0.39             | 1.33   | 0.02       | n.a.   | 0.02    | n.a.       |
| TiO₂      | 0.04      | 0.72   | 3.81             | 0.38   | 0.54       | n.a.   | 0.01    | n.a.       |
| Cr₂O₃     | 0.02      | 0.02   | 0.07             | 0.02   | 2.63       | n.a.   | 0.08    | n.a.       |
| MnO       | 0.04      | 0.01   | 0.01             | 0.07   | 1.84       | n.a.   | 0.60    | n.a.       |
| NiO       | b.d.l.    | b.d.l. | b.d.l.           | 0.01   | 0.15       | n.a.   | 0.12    | n.a.       |
| Total     | 101.09    | 101.75 | 100.72           | 100.84 | 99.74      | 95.5   | 98.86   | 99.02      |

**Table 2.** Intended phase compositions of the raw mixtures KMO1 and KMO2 (wt.%).

|          | C₂S | C₄A₃S | C₄AF | Total |
|----------|-----|-------|------|-------|
| KMO1     | 65  | 20    | 10   | 95    |
| KMO2     | 50  | 35    | 10   | 95    |

**Table 3.** Raw mix design (wt.%).

| Raw Mix | Limestone | Gypsum | Bauxite | Flysch | Steel Slag | Quartz | Calcite | Mill Scale | Total |
|---------|-----------|--------|---------|--------|------------|--------|---------|------------|-------|
| KMO1    | 40.3      | 4.2    | 5.1     | 38.9   | 7.7        | -      | 3.8     | -          | 100.0 |
| KMO2    | 44.1      | 7.2    | 13.7    | 13.9   | 11.1       | 4.1    | 5.6     | 0.3        | 100.0 |

Cements were prepared from clinkers sintered at 1250 °C, as this was the lowest temperature at which the targeted phase composition was reached. Cement clinkers and white titanogypsum were ground using a laboratory disc mill to pass through a 125 μm sieve. The proportion of gypsum added to the clinkers depended on the clinker phase composition intended. Gypsum was blended with the ground clinkers at a gypsum/calcium sulfoaluminate molar ratio of 1 [27]; 5.60 wt.% in the KMO1 sample and 9.90 wt.% in KMO2. The prepared cements were homogenized in a Turbula for 1.5 h and further milled to a similar Blaine specific surface area (SSA) of approximately 4900 ± 150 cm²/g in order to eliminate the influence of the particle size on the reactivity of cement (Table 4). However, despite the uniform Blaine SSA, the measurement of the particle size distribution (PSD) indicated that the fineness of the cement was greater in the KMO2 sample compared to KMO1 (Table 4, Figure 1).
Table 4. Specific density and Blaine specific surface area (SSA) of the belite-sulfoaluminate (BCSA) cements from raw mixtures KMO1 and KMO2 sintered at 1250 °C (Blaine method, ToniPERM Standard Model 6578, ToniTechnic by Zwick, Ulm, Germany).

| Cement  | KMO1 1250 °C | KMO2 1250 °C |
|---------|--------------|--------------|
| Specific density (g/cm³) | 3.17 | 3.03 |
| Blaine SSA (cm²/g) | 4890 | 4930 |

Figure 1. Particle size distribution (PSD) of cements from raw mixtures KMO1 and KMO2 sintered at 1250 °C, as determined by laser diffraction (Microtrac SYNC Model 5001, Microtrac Retsch GmbH, Haan, Germany; dry operation).

2.2. Methods

The phase composition of the cement clinkers was determined using a PANalytical Empyrean X-ray diffractometer (Malvern Panalytical, Malvern, UK) equipped with CuKα radiation and a PIXcel 1D detector. The samples were ground to a particle size of less than 63 µm. Samples were manually backloaded into a circular sample holder with a 10 mm diameter in order to mitigate the possible preferred orientation. The samples were measured at 45 kV at a current of 40 mA, in the range of 4 to 70 ° 2θ, at a step size of 0.026° 2θ with a scan step time of 197 s. Analysis of X-ray diffraction patterns was performed using XPert High Score Plus diffraction software v. 4.9 from PANalytical, using PANICSD v. 3.4 powder diffraction files. The Rietveld refinement method was conducted using crystal structures by Snellings et al. [28]. The structures for cubic and orthorhombic calcium sulfoaluminate were adopted from the work of Cuesta et al. [29,30].

SEM/EDS analysis was performed to study the morphology, size, and distribution of the phases, as well as to assess which minor elements were incorporated into the phases of the clinker. Uncoated polished cross sections of clinker pellets in vertical profiles were examined by a JEOL IT500 LV Scanning Electron Microscope (SEM) equipped with an Energy Dispersive X ray spectrometer (EDS) with a W filament, operated at an accelerating voltage of 15 kV in low vacuum mode at a working distance of 10 mm.

The hydraulic reactivity of cements prepared from clinkers sintered at 1250 °C was assessed by isothermal conduction calorimetry using a TAM Air calorimeter (TA Instruments). In total, 12 g of cement (ground to below 0.125 mm) with a water/cement ratio of 0.5 was mixed with an Ultra TurraX tube dispenser for 1 min. Then, 6 g of mixed paste was placed into glass vials and immediately put into the calorimeter. The heat evolution of cement was evaluated for 7 days at 20 °C.

To determine the compressive strength of cement prepared from clinkers sintered at 1250 °C, cement pastes with a water/cement ratio (w/c) of 0.5 were poured into prismatic molds with dimensions of 10 × 10 × 25 mm. At 24 h, the cement paste samples were demolded and cured in a humidity chamber at a temperature of 20 ± 1 °C and 95% relative humidity until they were tested under laboratory conditions. The compressive strength of cement pastes was determined at 7 and 28 days, using a ToniNORM test device.
(ToniTechnic by Zwick) at a load rate of 0.05 kN/s. The results are given as the average of four measurements for each cement mixture.

3. Results
3.1. Clinkers
3.1.1. X-ray Powder Diffraction

The phase composition of cement clinkers KMO1 and KMO2, sintered at different temperatures, was determined by Rietveld refinement (Table 5). In both types of clinker, the overall content of the main clinker phases was closest to the targeted values following sintering at 1250 °C (Table 5). The most abundant phase was C2S, which was mainly presented in beta form. During cooling, less than 1 wt.% of the beta C2S was transformed into gamma modification in both KMO1 and KMO2 (Table 5). The amount of belite did not significantly vary between clinkers sintered at 1200, 1250, and 1300 °C (Table 5). The second most abundant phase was C4A3S, which presented in orthorhombic and cubic form. The orthorhombic modification dominated in both clinkers at all sintering temperatures (Table 5). The highest amount of C4A3S developed at 1250 °C in both KMO1 and KMO2 clinkers (Table 5).

Table 5. Phase composition of cement clinkers KMO1 and KMO2, sintered at different temperatures, as determined by Rietveld refinement (wt. %). Rwp presents the weighted profile R-factor and GoF represents the goodness of fit.

|             | KMO1     | KMO2     |
|-------------|----------|----------|
| C4A3S - orthorhombic | 11.5     | 20.5     |
| C4A3S - cubic        | 4.2      | 15.1     |
| Σ C4A3S              | 15.7     | 35.6     |
| C2S-beta             | 64.6     | 64.5     |
| C2S-gamma            | 0.1      | 0.7      |
| Σ C2S                | 64.7     | 51.9     |
| C4AF                 | 9.9      | 2.2      |
| Perovskite           | 1.1      | 5.2      |
| Mayenite             | 5.3      | 3.3      |
| Periclase            | 2.4      | 2.3      |
| Aphthitalite         | 0.6      | -        |
| Langbeinite          | 0.3      | -        |
| Arcanite             | -        | -        |
| Total                | 100.0    | 100.0    |
| Rwp                  | 2.2      | 2.7      |
| GoF                  | 2.2      | 2.7      |

With regards to the C4AF phase, the targeted amount was reached at all three sintering temperatures in KMO1, while the amount of C4AF detected in KMO2 was much lower than the value predicted, only reaching around 2 wt.% (Table 5). The amount of ferrite remained constant between 1250 and 1300 °C, and then increased slightly between 1200 and 1250 °C (Table 5). However, perovskite was present in both clinkers, with less than 1 wt.% detected in KMO1, compared to more than 5 wt.% in KMO2 (Table 5). Ferrite is the artificial equivalent of natural mineral brownmillerite, having a structure similar to perovskite, which allows the substitution of various cations [31]. However, Ti was mainly incorporated into the raw clinker mixtures as a result of the presence of bauxite (Table 1), meaning the higher amount of perovskite in KMO2 can be explained by the larger amount of bauxite used in the KMO2 mixture (Table 3).

Among the minor phases, mayenite and periclase were detected in both clinkers. The amount of mayenite was higher in the KMO2 sample due to an increased content of the calcium sulfoaluminate phase. In both KMO1 and KMO2, the highest amount of mayenite was detected in samples sintered at 1200 °C, when the C4A3S was forming, while the lowest amount of C4A3S occurred at 1250 °C, which corresponds to the optimal
sintering temperature at which the targeted values were reached. The periclase (free magnesia (f-MgO)) content was similar for KMO1 and KMO2, with values remaining constant across all sintering temperatures (Table 5). The amount of periclase, determined by Rietveld refinement, was slightly enhanced in the two clinkers. For BCSA clinkers synthesized by including either bottom ash or fly ash, values are reported to be between 0.4 and 2.3 wt.% [16,32,33]. The main source of Mg was steel slag (Table 1); however, the overall amount of MgO introduced in both raw clinker mixtures was around 2 wt.%. The values are below the limit, if compared to the OPC clinker requirements, where the content of MgO should not exceed 5 wt.% [34] in order to avoid the formation of dead-burned MgO with a very slow hydration rate, as the late expansion of MgO would jeopardize the cement soundness [35]. At all sintering temperatures, some aphthitalite (K$_3$Na(SO$_4$)$_2$), langbeinite (K$_2$Ca$_2$(SO$_4$)$_3$), and arcanite (K$_2$SO$_4$) were also detected in KMO1 (Table 5), as a consequence of the higher content of fly ash, which contains alkalis.

3.1.2. SEM/EDS

Clinkers showed a clear decrease in porosity as the sintering temperature increased (Figure 2). In both mixtures, a temperature of 1200 °C was sufficient for developing the main reactive phases. However, X-ray Powder Diffraction (XRD) analysis revealed that the targeted amounts were not achieved at 1200 °C. Subhedral to euhedral C$_2$S and C$_4$A$_3$S grains were already clearly developed at 1200 °C in both KMO1 and KMO2 clinkers. However, at 1200 °C, the clinkers showed an inhomogeneous structure, as C$_4$AF was not present in the interstitial areas, but formed in individual nests (Figure 2). In both mixtures, sintering at 1250 °C resulted in a homogeneous structure, formed by subhedral to euhedral C$_2$S and C$_4$A$_3$S grains and C$_4$AF in the interstitial area (Figure 2). Raising the sintering temperature to 1300 °C led to structure densification, and increased the number of euhedral C$_2$S and C$_4$A$_3$S grains (Figure 2). Figure 3 shows elemental EDS maps for the KMO1 and KMO2 clinkers sintered at 1200, 1250, and 1300 °C.

Furthermore, the amounts of minor individual elements measured (EDS point measurements and SEM statistics), incorporated in the main clinker phases, are presented as sequences, in order of the maximum to minimum quantity detected. Belite enables various element substitutions [12,23,36–38]. It exhibited Al, Mg, Fe, S, K, Cr, Ti, Na, Ni, and Mn in the KMO1 clinker and Al, S, Fe, Mg, Cr, Ti, K, Na, Ni, and Mn in the KMO2 clinker. The higher amount of S detected in KMO2 corresponds to a higher gypsum content (Table 3). On the other hand, the higher Mg content of the KMO1 belites, as detected by EDS analysis, cannot be explained by a higher input from the raw materials (Tables 1 and 3). However, the highest amount of Mg in KMO1 belites was detected at 1200 °C, where the sintering was not complete, as evident from the EDS maps (Figure 4a). In both clinker mixtures, the highest amount of Al was incorporated into belite at 1250 °C (Figure 4a,b). On the other hand, the amount of Mg incorporated varied between the clinker mixtures. In KMO1, the amount of Mg in the belite phase clearly decreased with an increase in the sintering temperature (Figure 4a), while in KMO2 belites, the maximum amount of Mg was detected at 1250 °C (Figure 4b). In both clinkers, belites included the maximum amount of S at 1250 °C. The amount of Fe in the KMO1 belites decreased with a rise in the sintering temperature (Figure 4a), while in KMO2, the maximum amount of Fe incorporated in belite occurred at 1250 °C (Figure 4b). The maximum amount of K incorporated into belites occurred at 1200 °C in KMO1, while in KMO2, the K content of belites did not change as a result of the sintering temperature.

In both clinkers, calcium sulfoaluminate included minor elements in the same sequence, with Si, Fe, and Mg, followed by K, Cr, Ti, Na, Mn, and Ni. The quantity of Si decreased with rising temperatures in both KMO1 and KMO2, while the maximum amount of Fe and Mg occurred at 1250 °C (Figure 4c,d). The high Si content can be attributed to the fact that calcium sulfoaluminate is an end member of sodalite solid solution, in which Si is replaced by Al [39,40]. Consequently, increasing the sintering temperature resulted in decreased amounts of Si (Figure 4c,d).
Figure 2. The phase compositions of clinkers KMO1 (a) and KMO2 (b), sintered at 1200, 1250, and 1300 °C and shown at different magnifications (Scanning Electron Microscope (SEM)/ Backscattered-Electron (BSE) microphotographs). Clinker phase designations: C$_2$S = belite; C$_4$A$_3$S = calcium sulfoaluminate; and C$_4$AF = ferrite.
Figure 3. Elemental Energy Dispersive X ray spectrometer (EDS) maps for KMO1 (a–c) and KMO2 (d–f), sintered at 1200 °C (a,d), 1250 °C (b,e), and 1300 °C (c,f).

In KMO1, ferrite included Si, Ti, Mg, Mn, Ni, Cr, S, K, and Na, which slightly differed to the sequence of Si, Ti, Mg, Ni, Mn, S, Cr, K, and Na incorporated in KMO2. In both KMO1 and KMO2, the maximum amount of Si was incorporated into the ferrite phase at 1250 °C (Figure 4e,f). However, it has already been reported that Si and S commonly replace Fe or Al in a ferrite structure [41]. In both clinkers, the amount of Ti incorporated into the ferrite phase increased when raising the sintering temperature (Figure 4e,f). In KMO1, the maximum amount of Mg was incorporated into ferrites at 1250 °C, while in KMO2, the amount of Mg was constant across all sintering temperatures. In both clinkers, the maximum amounts of Ni and Cr were incorporated into ferrite at 1200 °C (Figure 4e,f).

Amongst the minor clinker phases, EDS mapping identified Mg-rich areas of periclase, which were seen in both clinker mixtures across all three sintering temperatures (Figure 3). Although steel slag contains a large amount of MgO (Table 1), the overall amount of Mg in clinkers was not substantially increased as clinker raw mixtures only incorporated around 8 or 11 wt.% of steel slag (Table 3). Moreover, the amount of periclase, determined by Rietveld refinement, was not very considerable (Table 5). The Ti-enriched areas correspond to perovskite, which was also identified in all clinkers (Figure 3a–d), but it was present to a greater degree in the KMO2 mixture (Figure 3d–f), as also supported by the XRD results (Table 5). Ca- and Al-enriched interstitial areas were observed in all clinkers, which can be attributed to the mayenite phase (Figure 3). KMO1 clinker mixtures also showed some K- and S-enriched areas of K-sulphate minor phases (Figure 3a–c).
The incorporation of minor elements into the main clinker phases of the KMO1 and KMO2 clinkers sintered at 1200, 1250, and 1300 °C, as determined by EDS analysis (wt.%): (a) Belite (C₂S) in KMO1; (b) C₂S in KMO2; (c) calcium sulfoaluminate (C₄A₃S) in KMO1; (d) C₄A₃S in KMO2; (e) ferrite (C₄AF) in KMO1; and (f) C₄AF in KMO2. The number of grains measured is given below the temperature in each individual diagram.

3.2. Cements
3.2.1. Isothermal Conduction Calorimetry

The hydration heat of the cements analysed is shown in Figure 5. The results showed that the kinetics of the two cements differ significantly.

Due to the external mixing method used, the first peak assigned to the heat of wetting and the early reaction during hydration [27,42,43] was not included. As can be seen from the results, the KMO1 cement reacted faster than the KMO2. Namely, the induction period for the KMO1 was very short following the first peak, lasting up to about 2 h of hydration in comparison to KMO2, where the induction period lasted until about 6 h of hydration.
before the main peak occurred. This period is attributed to the dissolution of clinker and the early formation of ettringite [27,43,44]. As follows, the main peak then occurred, which is attributed to the setting and hardening of the paste and the dissolution of calcium sulfoaluminate, together with gypsum, and the formation of ettringite and aluminium hydroxide [42,45,46]. This peak occurred after 4 h in KMO 1 and after 7 h in KMO2. Despite the fact that KMO2 contained a higher amount of highly reactive calcium sulfoaluminate phase, and that it had slightly finer particles compared to the KMO1, according to PSD measurement of the cement, which enhances its reactivity [47], the KMO1 reacted faster. However, this could be due to the presence of alkali sulfates (arcanite, aphthitalite, and Ca-langebeinite) in the KMO1. As already reported elsewhere, alkali sulfates have a significant effect on the hydration rate and setting time of cement. Namely, according to data in the literature [48–50], the reactivity of cement is accelerated in the presence of alkali sulfates. In addition, the KMO1 sample in comparison to KMO2 contains more ferrite, which is an active hydraulic phase, while it contains less perovskite, which is an inert phase [28–30]. Furthermore, the KMO2 sample displayed a second heat maximum after 11.5 h of hydration, which was attributed to the depletion of gypsum and the subsequent precipitation of monosulfate [42,43,51]. In KMO1, on the other hand, where a higher calcium sulfoaluminate amount was present, only a shoulder was observed at 6.5 h of hydration. This was probably due to the lower gypsum content in this mixture, which favored the formation of ettringite over monosulfate, and the peak was shifted to an earlier time [42,52].

![Hydration heat flow of the KMO cements.](image)

The cumulative heat results are shown in Figure 6 and Table 6. After 7 days of hydration, the amount of cumulative heat released was smaller in the KMO1 cement (176.6 J/g) than in KMO2 (223.8 J/g), indicating that more heat was released during the hydration of KMO2. This was attributed to a higher amount of calcium sulfoaluminate and a higher degree of hydration in the KMO2 cement.

3.2.2. Compressive Strength

As can be seen from the results in Table 7, the compressive strengths of the cement pastes vary significantly between 7 and 28 days of hydration, the extent of which depends on the cement mixture (KMO1 or KMO2). The compressive strength was higher in the KMO2 cement than the KMO1 cement at both 7 and 28 days, which is mainly related to
the increased calcium sulfoaluminate content (35 wt.% of C\(_4\)A\(_3\)S in KMO2 compared to 20 wt.% of C\(_4\)A\(_3\)S in KMO1).

Figure 5: Hydration heat flow of the KMO cements.

The cumulative heat results are shown in Figure 6 and Table 6. After 7 days of hydration, the amount of cumulative heat released was smaller in the KMO1 cement (176.6 J/g) than in KMO2 (223.8 J/g), indicating that more heat was released during the hydration of KMO2. This was attributed to a higher amount of calcium sulfoaluminate and a higher degree of hydration in the KMO2 cement.

Figure 6. Cumulative heat of hydration in KMO cements for up to 7 days of hydration.

Table 6. Cumulative heat of the KMO cements at 7 days of hydration.

|          | KMO1 | KMO2 |
|----------|------|------|
| Cumulative heat (J/g) | 176.6 | 223.8 |

Table 7. Compressive strength of the KMO cements.

|          | KMO1 | KMO2 |
|----------|------|------|
| Compressive strength, 7 days (N/mm\(^2\)) | 17.3 ± 1.2 | 24.6 ± 1.2 |
| Compressive strength, 28 days (N/mm\(^2\)) | 21.1 ± 1.2 | 33.2 ± 0.6 |

Consequently, more gypsum was needed in the KMO2 cement mixture for obtaining an equal molar ratio. Ettringite and aluminum hydroxide are formed from calcium sulfoaluminate in the presence of water and a source of sulphate. The more calcium sulfoaluminate is added, the more ettringite forms, the higher the volume of hydrates, and the lower the amount of available water and available space for growing hydrates [26,51,53]. It is ettringite which gives the initial strength (reaction detected by isothermal calorimetry). Since more belite reacts during later age periods, this will further contribute to the long-term development of compressive strength [54]. The increase in the compressive strength of the cements from 7 to 28 days is smaller in KMO1 than in the KMO2 cement. Both studied cement clinkers contained slightly increased periclase (f-MgO) contents. The formation of Mg(OH)\(_2\) during f-MgO hydration leads to inner stresses, causing a reduction in the compressive strength [55]. Nevertheless, generally, as observed for OPC, there was a decrease in the compressive strength with increased f-MgO contents (> 3 wt.%), especially at later ages [55–57]. At the same time, there was a compressive strength reduction with an increased f-MgO content, relative to a reference paste without additional f-MgO (ordinary Portland cement mix), mainly attributed to the increase in porosity and lower hydration of f-MgO mixtures compared to MgO-free mixtures [58].

4. Conclusions

The present study presents the results of an investigation into the use of steel slag in belite-sulfoaluminate cement clinkers.

For the two clinker phase compositions targeted, it was possible to incorporate between 7.7 and 11.1 wt.% of the steel slag into the raw clinker meal. The results showed
that it was possible to achieve the targeted phase composition of both types of clinker at a sintering temperature of 1250 °C. In addition to the main phases of belite, calcium sulfoaluminate, and ferrite, minor phases were also identified. While mayenite, perovskite, and periclase were identified in both clinkers at all temperatures, alkali sulfates were mainly identified in the KMO1 clinker, since it contained more fly ash with enhanced alkalis. Incorporation of the steel slag especially resulted in the formation of periclase in the two clinkers, since it contains elevated levels of MgO. Furthermore, a higher content of Ti-bearing bauxite in the KMO2 clinker raw mixture promoted the formation of perovskite. A higher amount of mayenite was observed at the lowest sintering temperature (1200 °C) in both clinker mixtures, but it was higher in KMO2 due to a higher amount of the calcium sulfoaluminate phase. As evidenced by SEM/EDS, a clear porosity decrease and clearer phase development were seen in the clinkers with increased sintering temperatures. Furthermore, various foreign ions were incorporated into the main phases.

The hydration kinetics of the cements prepared by clinkers were influenced by both the major and minor phases. The heat of hydration was higher in the KMO2, which was made from clinker with a higher content of very reactive calcium sulfoaluminate. Similarly, the compressive strength of cement was higher in the KMO2 cement at both 7 and 28 days. The latter also showed a greater increase in compressive strength development compared to KMO1, with a larger proportion of belite.

The results confirmed that steel slag can be incorporated into belite-sulfoaluminate cement clinkers. The cements showed appropriate hydraulic characteristics and developed a sufficient compressive strength.

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