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## 1 ALKALI-ACTIVATED BINDERS BASED ON GROUND GRANULATED BLAST

## 2 FURNACE SLAG AND PHOSPHOGYPSUM

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## 18 Abstract

19	The effect of phosphogypsum (PG) in alkali-activated ground granulated blast furnace slag (GGBFS)
20	on the reactivity, strength development and final matrix properties was investigated, as a function of
21	alkali activator. The results were compared with alkali-activated reference samples from GGBFS. PG
22	completely dissolves and takes part in solid reaction product formation. Upon PG incorporation,
23	portlandite and ettringite are initially formed after dissolution. The hardened binder consists mainly
24	of amorphous hydration products, intermixed with thenardite and minor amounts of secondary
25	gypsum and merwinite. PG incorporation resulted in faster initial and delayed final setting time,
26	while it enhanced the compressive strength and amorphous phase development when sodium
27	hydroxide was used. PG gave rise to a decrease of Al/Si and Ca/Si ratios in the C-A-S-H gels and a
28	higher polymerized network.
29	
30	Keywords
31	Super-sulphated cement, alkali-activated binder, ground granulated blast furnace slag,
32	phosphogypsum, hydration, building material
33	
34	1. Introduction
35	The search for new binders, wherein Ordinary Portland Cement (OPC) gets partly or totally replaced
36	by other resources, can contribute to the reduction of energy usage and carbon dioxide generation
37	caused by OPC's manufacture. Super-sulphated cements (SSC) are alternative binders which are free
38	(or almost free) of OPC, and consist in most cases of 70-90% ground granulated blast furnace slag
39	(GGBFS), 10-20% gypsum and an alkali activator [1–3]. Next to their environmental benefits [4,5], SSC
40	show promising potential for industrial applications because of their technical performance, e.g. high

41 strength [6] and excellent resistance to chemically aggressive surroundings [1,3,7].

42 Considering the availability of various gypsum-rich waste streams, there is a strong interest in finding 43 applications that could in effect be both economically viable and sustainable. In this respect, phosphogypsum (PG) forms a potential candidate as gypsum source in the synthesis of SSC. PG is a 44 45 by-product originating from the production process of phosphoric acid, consisting mainly of 46 CaSO<sub>4</sub>.2H<sub>2</sub>O and impurities, such as fluoride, remnants of phosphoric acid, heavy metals (e.g. Cd or Cr) and naturally occurring radionuclides (mainly from the <sup>238</sup>U decay chain). Worldwide, large 47 48 quantities are disposed in ponds or heaps without purification, because existing purification 49 techniques are not effective [8,9] and this practice may increase the costs [10]. For this reason, PG 50 has very low recycling rates and it is predicted that the total amount disposed will range from 7 to 8 51 billion tons by 2025 [11], requiring large disposal areas and necessitating proper management to 52 avoid environmental pollution. Furthermore, a future increase in production volumes is expected 53 because of increasing food production demand accompanied by increasing fertilizer production 54 [9,11,12]. Because of these large volumes, the construction industry is considered to form a possible 55 valorization route. However, if PG is considered for reuse in building applications, the maximum level 56 of naturally occurring radionuclides in building materials is being regulated by the European Basic 57 Safety Standards (EU-BSS) [13], provoking limitations. Therefore, this work is complementary to a 58 previous investigation that deals with the radiological impact of the same binders [14], where it was 59 concluded that maximum 10 wt% PG could be incorporated in order to produce innovative, 60 environmental-safe building materials, which integrate industrial by-products yet comply with 61 existing safety standards.

Slags from iron and steel manufacture, such as GGBFS and ladle slag, respectively, can even react
with gypsum without an activator, due to their alkalinity (i.e. high CaO + MgO content) [1]. Despite,
fluoride and remnants of phosphoric acid in PG decline the reaction rate and consequently increase
the setting time [9]. To achieve a sufficient reactivity, an alkaline surrounding is preferable, provided
by an alkaline solution in this investigation.

67 SSC from GGBFS, PG and an alkali activator could form a new class of alternative sustainable binders, by avoiding the use of OPC and creating a valorization route for PG. However, due to its impurities, 68 the effect of PG incorporation on the reactivity, strength development and final matrix properties 69 70 should be investigated, which forms the purpose of this study. Also, the use of different alkali 71 activators is investigated. Reactivity is assessed by recording the setting time of the pastes, while 72 isothermal calorimetry and in-situ X-ray diffraction (XRD) analysis were employed to monitor the 73 reaction rate of the product formation. Final matrix properties were investigated by measuring 74 compressive strength and applying micro-chemical analysis. Fourier-transformed infrared (FTIR) 75 spectroscopy and XRD were used for investigation of the hydration products.

76

## 77 2. Materials and methods

## 78 2.1 Precursors

79 GGBFS was derived from iron extraction in a Belgian steel producing company. PG originated from a PG processing plant in Gdansk (Poland) and was provided by the International Atomic Energy Agency 80 (IAEA) (reference material n° 434) [15]. After deriving, GGBFS was dried at 110 °C for 24 h and milled 81 82 to a Blaine value of 4050 ± 200 cm<sup>2</sup>/g, calculated according to EN 196-6 [16]. The density of GGBFS 83 was measured with a Quantachrome Multipycnometer (MPV-6DC) and found to be 2.9 g/cm<sup>3</sup>, 84 according to ASTM C204 [17]. The chemical composition of GGBFS was analyzed by X-ray 85 fluorescence analysis (XRF) using a Philips PW 1830 instrument, and is summarized in Table 1. PG was 86 used as received with a  $d_{50}$  value of around 3  $\mu$ m and the matrix composition was (in wt%): 96.0 87 CaSO<sub>4</sub>.2H<sub>2</sub>O, 1-2 P<sub>2</sub>O<sub>5</sub>, 1.2 F<sup>-</sup>, 1.0 SiO<sub>2</sub> and 0.2 Al<sub>2</sub>O<sub>3</sub> [15]. FTIR spectroscopy (Bruker Vertex 70) was performed on GGBFS, PG and a GGBFS/PG mixture by proportion of 9/1 (which is the applied 88 89 precursor weight ratio, see Table 2). Spectra were acquired from 4000 cm<sup>-1</sup> to 450 cm<sup>-1</sup>, at a 90 resolution of 4 cm<sup>-1</sup> and with 32 scans per measurement, supported by Opus software. Prior to 91 measurement, a background spectrum was recorded. XRD measurements were performed on GGBFS

92 and PG. They were milled for 5 min in a McCrone micronizing mill with corundum grinding elements and ethanol to obtain accurate fineness. As an internal standard material, 10 wt% of analytical-grade 93 crystalline ZnO (purity 99.9%, Merck) was added. An automated diffractometer with Cu-Kα-radiation 94 95 (D2 PHASER, Bruker) with a Lynx-eye super speed position sensitive detector was applied for the XRD 96 analysis. An anti-scatter slit positioned 3 mm above the sample was used to protect the detector 97 from abundant X-rays at lower 2 $\theta$  degrees. In order to reduce preferred orientation, the samples were prepared using the back loading technique. The settings for the X-ray tube generator were 30 98 99 kV and 10 mA. To improve statistics, the samples were rotated at 15 rpm during the measurement. 100 Diffractograms were recorded in continuous PSD fast mode between 5° and 70° 20 at 0.02° step size 101 and a counting time of 0.3 s per step. The diffractograms were qualitatively investigated with EVA 102 V.3.1 (Bruker AXS). Quantitative analysis was performed with MAUD [18] based on the Rietveld 103 method [19–21]. From the known initial ZnO content of 10 wt%, the crystalline and amorphous 104 phase contents were recalculated.

105

Table 1: Chemical composition of GGBFS

Chemical compound	Wt%				
SiO <sub>2</sub>	36.2 ± 0.2				
CaO	40.3 ± 0.5				
Al <sub>2</sub> O <sub>3</sub>	$11.4 \pm 0.2$				
MgO	8.2 ± 0.1				
SO <sub>3</sub>	$1.1 \pm 0.1$				
TiO <sub>2</sub>	$0.8 \pm 0.1$				
Na <sub>2</sub> O	$0.8 \pm 0.1$				
K <sub>2</sub> O	0.5 ± 0.1				
Fe <sub>2</sub> O <sub>3</sub>	0.3 ± 0.1				

106

## 107 **2.2 Sample's mix design**

108 To investigate the effects of PG incorporation, six mixtures were designed. The alkali activators used

were the combination of a sodium silicate solution (molar ratio  $SiO_2/Na_2O = 3.3$  and 65% water,

supplied by ABCR GmbH), sodium hydroxide pellets (grade 99%, supplied by Chem-Lab) and distilled

111 water (ASTM type II), and were prepared one day prior to sample synthesis. The mix designs are

summarized in Table 2. The mixing time amounted 3 min. An alkali activator/precursor ratio of 0.6
was chosen because a decline in the workability was observed upon PG incorporation, caused by its
high specific surface area. This ratio was retained for the synthesis of pure GGBFS samples for the
sake of comparison.

116

Table 2: Sample's mix design (partly adapted from [14])

Sample	wt% GGBFS	wt% PG	Alkali activator/Precursor	SiO <sub>2</sub> /Na <sub>2</sub> O	$H_2O/Na_2O$
SS1	90	10	0.6	0.75	20.0
SH2	90	10	0.6	0	27.8
SH3	90	10	0.6	0	18.5
SS4	100	0	0.6	0.75	20.0
SH5	100	0	0.6	0	27.8
SH6	100	0	0.6	0	18.5

117

## 118 2.3 Physico-chemical analysis

119 Isothermal calorimetry was executed at 20 °C using a TAM-Air Isothermal Calorimeter (TA 120 Instruments). First the precursor was inserted in the admix ampoule and the alkali activator into the 121 syringe. Then the ampoule was added into the calorimeter and once thermal equilibrium was 122 obtained, precursors and alkali activators were mixed for 3 min prior to measurement. For 123 determination of the setting time, pastes were prepared and loaded in a Vicat apparatus (Matest 124 E044N Vicatronic) in accordance with EN 196-3 [22]. For selected samples, in-situ XRD (D2 PHASER 125 (Bruker)) was applied. The surface of the paste was smoothed and the sample holder was sealed with 126 Kapton film to prevent water loss and carbonation. The hydrating paste was measured for 22 h and 127 diffractograms between 6° and 55° 20 were assembled at time intervals of 13 min and with a step-128 size of 0.02° 20 and counting time of 0.3 s per step. An anti-scatter slit was positioned 1 mm above 129 the sample. The settings for the X-ray tube generator were 30 kV and 10 mA. The temperature inside 130 the diffractometer case was 20 °C during the measurements. The diffractograms were qualitatively 131 investigated with EVA V.3.1 (Bruker AXS). For compressive strength tests, fresh paste mixes were cast 132 in polymer coated steel molds (20 mm x 20 mm x 80 mm), wrapped in plastic foil to prevent drying

133 and cured at 20 °C. The samples were demolded after 24 h and further cured at 20 °C in sealed vessels. After 7 and 28 days, five specimens of each formulation were tested with an Instron 5985 134 machine equipped with a 250 kN load cell. Quantitative elemental point analysis was performed after 135 136 28 days of curing with electron probe micro-analysis (EPMA) (Jeol JXA-8530F (FEG)). Specimens 137 sampled from the middle of the cured samples were embedded in epoxy resin, polished and coated 138 with a 25 nm C layer. The settings for the microprobe were 15 kV and 15 nA. The standards of 139 obsidian (for Al, Na and Si), apatite (for Ca and P), celestite (for S), pyrite (for Fe), periclase (for Mg) 140 and kaersutite (for K) were used. Also the background was measured under these conditions. 141 Mappings for Al, Ca, Mg, Na, S and Si were obtained with a dwell time of 40 ms per pixel in a mapped 142 area of 480 x 345 μm (pixel size 1 μm). FTIR spectroscopy was performed after 28 days of curing. 143 After their curing period, samples were dried at 40 °C till constant weight and subsequently crushed 144 in a porcelain mortar and passed through a 45 µm sieve. Measurements were carried out with the 145 same procedure as described earlier (section 2.1). XRD measurements were performed after 28 days 146 of curing. After their curing period, samples were dried at 40 °C in a laboratory oven till constant 147 weight whereafter they were grinded in a porcelain mortar. Subsequent milling, measurements and 148 analyses were performed as explained in section 2.1.

149

## 150 3. Results and discussion

## 151 **3.1 Isothermal calorimetry and setting time**

Fig. 1 shows the heat evolution curves, calculated based on total sample mass (precursor + alkaliactivator). Time zero represents the time after 3 min of mixing.





#### Figure 1: Heat flow

156 As GGBFS is used as precursor, the main reaction product formed upon alkali activation is expected 157 to be a calcium aluminosilicate hydrate (C-A-S-H) type gel [23], which is different from the calcium 158 silicate hydrate (C-S-H) generated in OPC hydration. Generally, C-A-S-H could be viewed as a C-S-H 159 wherein  $AI^{3+}$  is located within the silicate chains, more specifically at the tetrahedral bridging sites [23,24]. The incorporation of 10 wt% PG forms a source of  $Ca^{2+}$  and  $SO_4^{2-}$  ions enhancing the 160 161 formation of secondary reaction products. The rate of reaction product formation is inherently determined by various parameters, and consequently these reactions can occur simultaneously or in 162 sequence. The heat flow curves can be interpreted as the summation of heat released and consumed 163 164 by these reactions. For samples SS1, SH2 and SH3, an initial very intense peak with short duration 165 (from approximately 0.1 to 0.5 h) is observed, which corresponds to the formation of ettringite 166 (confirmed by in-situ XRD, section 3.3), being a hexacalcium aluminate trisulphate hydrate mineral, 167 with as chemical formula  $Ca_6Al_2(SO_4)_3(OH)_{12}$ . 26H<sub>2</sub>O [25]. In cement hydration, it was also found that 168 ettringite immediately forms after dissolution [26]. During its formation, crystalline water and a 169 framework of hydrogen bonds are formed upon consuming water from the alkali activator. After this, 170 a dormant period is observed, meaning that the formation of other reaction products is postponed, 171 most likely because of a reduced amount of water available for reactions and surface blockage of 172 GGBFS grains by ettringite, which initially hindered their further hydration. The onset of the peak of

173 reaction product formation is therefore shifted to later time and this peak is much broader as for pure GGBFS samples, indicating a slower reaction rate as a consequence of fluoride and remnants of 174 phosphoric acid in PG. The presence of silicates in the alkali activator, and a lower H<sub>2</sub>O/Na<sub>2</sub>O for 175 176 sodium hydroxide alkali activators, causes a shift of the peak of reaction product formation to earlier 177 times. The height of both the peak of ettringite formation (9.4 mW/g, 7.7 mW/g and 14.4 mW/g after 178 0.2 h for sample SS1, SH2 and SH3, respectively) and the peak during reaction product formation (2.3 179 mW/g at 1.8 h for SS1, 2.2 mW/g at 3.4 h for SH2 and 3.2 mW/g at 2 h for SH3) is inversely 180 proportional to the H<sub>2</sub>O/Na<sub>2</sub>O ratio of the alkali activator. The heat release during reaction product 181 formation is comparable with values found in literature on SSC [1]. 182 The curve shape of pure GGBFS-based binders are similar to data published elsewhere [27–29]. For 183 sample SH5, a dissolution peak is observed at first, whereafter a dormant period follows. 184 Subsequently, a broader exothermic peak is recorded. During the dissolution and the dormant 185 period, the concentration of dissolved species gradually increases, whereupon C-A-S-H formation 186 starts when concentrations reach a threshold [30,31]. For samples SS4 and SH6, there is no dormant 187 period, indicating that immediately after dissolution the concentration threshold was reached to 188 start C-A-S-H formation. The height of the peak during C-A-S-H formation is inversely proportional to 189 the H<sub>2</sub>O/Na<sub>2</sub>O of the alkali activator. Sample SS4 is characterized by a maximum heat flow of 2.9 mW/g after 0.3 h, for sample SH5 this was 2.7 mW/g after 1.2 h and for sample SH6 3.6 mW/g after 190 191 0.9 h. The presence of silicates in the alkali activator accelerates C-A-S-H formation, and a decrease 192 of H<sub>2</sub>O/Na<sub>2</sub>O for sodium hydroxide alkali activators results in a shift of the peak to earlier time. For 193 sample SS4, another broad peak after 5 h is observed, likely representing further densification of the 194 C-A-S-H structure.



## 196

#### Figure 2: Setting time

197 These thermal characteristics can be correlated to the initial and final setting times of the paste 198 samples, which are presented in Fig. 2. Time zero represents 3 min after the start of mixing. In 199 general, the incorporation of 10 wt% PG results in a faster initial setting (due to the high specific 200 surface area of PG and early ettringite formation) (i.e. a reduction of 0.1 h to 1.2 h) but slower final 201 setting (due to (1) a reduced amount of water available for the reactions, (2) initial surface coverage 202 of GGBFS grains by ettringite and (3) fluoride and remnants of phosphoric acid acting as set retarder 203 [32]) (i.e. an increase of 0.1 h to 2 h). The alkali activator with the highest  $H_2O/Na_2O$  (i.e. samples SH2 204 and SH5), led to highest initial and final setting times. The type and concentration of alkali activator 205 plays an essential role in order to control the setting times, which is already extensively reported in 206 literature on alkali-activated slags [33–35] confirming the results for samples SS4, SH5 and SH6. 207 Regarding the initial setting times, samples SS1, SH2 and SH3 fulfill the requirements defined in EN 208 15743 [2] on SSC (initial setting time ≥ 0.75 h), providing an appropriate time in real construction 209 applications.

210

## 211 3.2 In-situ XRD

212 In-situ XRD was performed for sample SH3 which displayed the highest heat flow peak from





232 which a certain amount of activator is necessary to develop a cascade of formation and dissolution 233 reactions. After about 12 h, the intensity of the peaks of portlandite and thenardite decreases 234 because the amorphous content is increasing. After 22 h, the hardened sample consists of 235 amorphous hydration products, intermixed with thenardite and portlandite. Because thenardite is a 236 salt, leftovers will give rise to high leaching rates of sodium and sulphate when in contact with water. 237 Therefore, ideally all the thenardite gets totally consumed during hydration. Also for samples SS1 and 238 SH2, thenardite was left after 28 days of curing (confirmed by XRD, section 3.6), which is not 239 desirable from the leaching point of view. On the contrary, the formation of thenardite is enviable 240 from the microstructural development point of view, giving rise to enhanced strength (section 3.3) 241 because of accelerated GGBFS dissolution.

242

## 243 **3.3 Compressive strength testing**

244 Fig. 4 shows the compressive strength (1 sigma) at 7 and 28 days. For all samples, an increase in 245 compressive strength is observed when extending the curing period, attributed to the development 246 of reaction products to different degrees as a function of sample's mix design. Compared with the 247 reference samples, the incorporation of 10 wt% PG for samples SS1, SH2 and SH3 resulted in strength 248 reduction (33.1% at 28 days) for the sodium silicate activated samples (SS1), while it caused a 249 strength gain for sodium hydroxide activated ones (SH2 and SH3) (55.3% and 89.0% at 28 days, 250 respectively). This is likely due to a slightly higher amorphous content (see further in section 3.6) of 251 SH2 and SH3, compared with SH5 and SH6, respectively. The strength reduction for SS1 agrees with a 252 lower amorphous content compared to SS4. Resulting from the presence of additional silicate 253 species, samples activated with sodium silicate (SS1 and SS4) showed significant higher strength 254 compared with sodium hydroxide activated samples, due to enhanced solid product formation and 255 the formation of a higher polymerized structure (see also FTIR, section 3.5). For sodium hydroxide 256 activated samples, an increase in Na<sub>2</sub>O concentration (SH3 and SH6) led to an increase in

compressive strength at 28 days (compared to SH2 and SH5, respectively), which can be explained by
a higher dissolution rate of precursors, resulting in more species available for reaction product
formation. The compressive strength at 28 days of pure GGBFS samples (SS4, SH5 and SH6) is
comparable with values found in literature, although the applied alkali activator/precursor ratio of
0.6 is higher than mostly used [38].



262



Figure 4: Compressive strength after 7 and 28 days of curing

264 Only for sample SS1, the compressive strength at 7 and 28 days exceeds the minimum legitimated in 265 the European standard for SSC [2]. Though, depending on the considered application, different 266 strengths are required, e.g. for driveways and footpaths typical compressive strengths are ranging 267 from 15 MPa to 25 MPa, while the strength for structural concrete for bridge deck slabs should 268 achieve 32 MPa. The compressive strength of superstructures ranges from 40 MPa to 50 MPa and 269 concrete pavements require 30 MPa [39]. Nonetheless, this is merely the first data and additional 270 testing at later ages, accelerated testing and repeating these tests on concrete samples is needed in 271 order to safely define proper applications.

272

## 273 3.4 Micro-chemical analysis

Micro-chemical analysis was performed after 28 days of curing and backscattered electron (BSE)
images (500 times magnification) are shown in Fig. 5. The samples demonstrate a heterogeneous
microstructure, consisting of a mix of solid reaction products (darkest areas) and undissolved GGBFS
particles (lightest areas). Upon PG incorporation, different reaction products are formed (indicated
by R1 and R2 in Fig. 5). R1 represents the C-A-S-H gel, while R2 could indicate secondary gypsum,
thenardite and/or monosulphate (see further in section 3.6). Microcracks presumably have
developed during sample preparation (when they are placed in vacuum for coating).



281

Figure 5: BSE images after 28 days of curing: a) SS1, b) SH2, c) SH3, d) SS4, e) SH5 and f) SH6 282 283 Quantitative elemental point analysis was performed, the results are not presented due to 284 unacceptable high uncertainties. It is not always straightforward to understand what is located 285 underneath the sample surface since the excited volume for such analysis lies in the micrometer 286 range, which is considerably large to elucidate a whole of intermixed solid hydration products. For 287 this reason, mappings are more reliable, which are shown in Fig. 6 for samples incorporated with PG. 288 Undissolved GGBFS particles are embedded in the C-A-S-H gel, while regions of secondary gypsum 289 (SG, see further in section 3.6) and thenardite (T) are clearly detectable.





296

Figure 6: Mappings of samples SS1, SH2 and SH3 after 28 days of curing

297

## 298 3.5 Fourier-transform infrared

299 FTIR spectra of the precursors (Fig. 7a) were recorded to visualize the change in the spectrum when 300 precursors get alkali-activated. For the sake of clarity, spectra were shifted upwards and have been 301 plotted from 3800 to 450 cm<sup>-1</sup>. GGBFS presented a stretching vibration (Si-O bonding) centred at 302 around 890 cm<sup>-1</sup>. The broadness of the stretching vibration of the GGBFS/PG mixture, centred at 303 around 900 cm<sup>-1</sup>, was larger than those for GGBFS, which is the reference to observe the hydration 304 process. For PG, the bands centred at 1105, 1090, 1005, 659 and 595 cm<sup>-1</sup> are assigned to the stretching and bending modes of sulphate. The bands at 1105 and 1090 cm<sup>-1</sup> correspond to the 305 306 asymmetrical  $v_3$  vibrations, the band at 1005 cm<sup>-1</sup> corresponds to the asymmetrical  $v_1$  vibrations, 307 and the ones at 659 and 595 cm<sup>-1</sup> to the asymmetrical  $v_4$  vibrations [40]. The stretching vibration of 308 the H<sub>2</sub>O molecules occurs at 3600, 3560 and 1616 cm<sup>-1</sup>. The bands at 675, 614 and 595 cm<sup>-1</sup> are 309 characteristic for anhydrite [41], indicating that some amount of CaSO<sub>4</sub>.2H<sub>2</sub>O was transformed to 310  $CaSO_4.0.5H_2O$  and  $CaSO_4$  (confirmed by XRD, section 3.6).



312

Figure 7: FTIR spectra of a) precursors and b) samples after 28 days of curing

313 As a result of the amorphous character of alkali-activated binders, disabling the applicability of XRD 314 for characterization of the non-crystalline phases, FTIR spectroscopy was applied to gain useful information. Fig. 7b shows the FTIR spectra of samples at 28 days. Since there were no major 315 changes within the 4000 to 1500 cm<sup>-1</sup> region, data were plotted from 1500 to 450 cm<sup>-1</sup> and the 316 spectra were shifted upwards. Because GGBFS was used as precursor, the formation of C-A-S-H gel as 317 318 main reaction product is expected [28,42–44]. The stretching vibration of Si-O-T (T stands for Al or Si) 319 units (generated by Q<sup>2</sup> units), commonly observed around 940 cm<sup>-1</sup> upon alkali activation [45–47], 320 provides valuable information about the constitution of reaction products. This stretching vibration is 321 related to a low-polymerized silicate network, wherein Si gets partly substituted by other elements 322 (e.g. Al), and where non-bridging oxygens (e.g. Si-O-Na) occur [48,49]. Other bands, which are also characteristic for the Si-O-T units, are the ones around 400-500 cm<sup>-1</sup> (deformation vibrations) and 323 around 700-800 cm<sup>-1</sup> (typical for Q<sup>1</sup> units) [50]. No Q<sup>3</sup> units were discovered (i.e. at 1200 cm<sup>-1</sup>). The 324 325 broader these Si-O-T bands, the more amorphous the material, while an increase in their 326 wavenumber designates a more polymerized network (enhanced Si content in the C-A-S-H gel). From 327 Fig. 7b, it can be seen that the strongest bands lay in the interval of 930-950 cm<sup>-1</sup>, and as stated 328 earlier, being the stretching vibration area for C-A-S-H gels from GGBFS activation [51]. The changes 329 located at the stretching vibration area for samples incorporated with 10 wt% PG (SS1, SH2 and SH3) 330 to slightly higher wavenumbers, could be attributed to the decrease of Al/Si and Ca/Si ratios in the

331 gels [52] and a higher polymerization network. On the contrary, sample SS4, activated with a sodium silicate solution, did not possess a change of the stretching vibration band to higher wavenumbers, 332 333 compared with sodium hydroxide activated ones (SH5 and SH6). An explanation for this behavior 334 could be that in spite of the fact that a higher polymerized structure was formed when using a 335 sodium silicate solution, the intensity of the absorption was diminished due to their slightly higher 336 amorphous nature (broader band), whereby the resulting peak was not shifted to higher energies. 337 On the other hand, this could also indicate that changing the SiO<sub>2</sub>/Na<sub>2</sub>O molar ratios of the alkali 338 activator has minor influence on the degree of polymerization. However, the first suggestion is 339 assumed to be more realistic regarding compressive strength (section 3.3) and confirmed by Q-XRD 340 (section 3.6). The stretching vibration band, and the ones located around 400-500 cm<sup>-1</sup> and 700-800 341 cm<sup>-1</sup>, show highest intensity for samples SH3 and SH6, most likely caused by an initially higher 342 precursor dissolution rate due to a higher Na<sub>2</sub>O content in the alkali activator. The band around 800 343 cm<sup>-1</sup> is less clear for samples activated with sodium silicate (SS1 and SS4), which is commonly 344 observed for silicates with high network modifiers [50]. All samples displayed OH<sup>-</sup> groups at around 345 1650 cm<sup>-1</sup> and 3600 cm<sup>-1</sup> (not presented in the figure), indicating the occurrence of chemically bound 346 water across the reaction products [53]. For sample SH6, carbonation has taken place, deduced from 347 the bands at 865 cm<sup>-1</sup> and 1444 cm<sup>-1</sup>, which are related to the asymmetric stretching vibration of O-348 C-O bonds [54,55]. Since carbonation was not observed for SH3, this phenomenon could be impeded by the incorporation of PG. The band observed at 1400 cm<sup>-1</sup> for all samples might indicate the 349 350 presence of hydroxyl groups in the material. The bands in the region from 500 to 700 cm<sup>-1</sup> may be 351 characteristic for iron-containing phases [56], but are in this case more probably caused by the Al-O 352 bonding. The bands at 1110 cm<sup>-1</sup> and 610 cm<sup>-1</sup> for samples SS1, SH2 and SH3 are nonexistent in the 353 spectra of alkali-activated binders containing exclusively GGBFS, and correspond to thenardite, 354 secondary gypsum and/or monosulphate (see further in section 3.6). The band at 1110 cm<sup>-1</sup> represents the asymmetric stretching vibration of SO<sub>4</sub><sup>2-</sup>, while the band at 610 cm<sup>-1</sup> corresponds to its 355 356 out-of-plane bending vibration [57,58].

363

## 358 3.6 XRD

Fig. 8a shows the diffractograms of the precursors. The asterisks indicate the peaks from the internal
ZnO standard. GGBFS was predominantly amorphous with a trace of merwinite (Ca<sub>3</sub>Mg(SiO<sub>4</sub>)<sub>2</sub>, 5.3
wt%). PG comprised a mixture of 33.2 wt% CaSO<sub>4</sub>.2H<sub>2</sub>O, 27.1 wt% CaSO<sub>4</sub>.0.5H<sub>2</sub>O, 32.1 wt% CaSO<sub>4</sub>
(also confirmed by FTIR, section 3.5) and 7.6 wt% amorphous phases.



Figure 8: XRD patterns of a) precursors and b) samples incorporated with PG after 28 days of curing 364 365 As a result of the highly amorphous character of alkali-activated binders, only little information about 366 the crystalline phases could be gained from their diffractograms after 28 days of curing (shown in Fig. 367 8b for samples incorporated with PG). The data were plotted from 5 to 40 20, the peaks from the 368 internal ZnO standard are marked with an asterisk. Peaks of gypsum, merwinite, thenardite and 369 calcite were detected. Carbonation (formation of calcite) had occurred during sample preparation 370 (wet milling with ethanol and subsequent evaporation), since it was not observed in their FTIR 371 spectra (section 3.5). Peaks of gypsum were not observed with in-situ XRD during the first 22 h. 372 However, the conversion of early ettringite results in the presence of secondary gypsum and 373 monosulphate (Ca<sub>4</sub>Al<sub>2</sub>SO<sub>4</sub>(OH)<sub>12</sub>.6H<sub>2</sub>O) according to Eq. 1 [37]:

374 
$$Ca_6Al_2(SO_4)_3(OH)_{12}$$
.  $26H_2O \rightarrow Ca_4Al_2SO_4(OH)_{12}$ .  $6H_2O + 2CaSO_42H_2O + 16H_2O$  (Eq. 1)

- 375 Despite, monosulphate is a poorly crystalline phase and hard to detect by XRD [59–61]. Portlandite
- 376 completely dissapeared after 28 days of curing.



378

#### Figure 9: Q-XRD of samples after 28 days of curing

379 In Fig. 9, the quantitative analysis of X-ray data is presented. From these data it is concluded that the 380 main reaction product was an amorphous C-A-S-H gel for all samples. The amorphous content of 381 sample SS1 was lower compared with SS4, while samples SH2 and SH3 showed increased amorphous 382 content compared to SH5 and SH6, respectively. When comparing Fig. 4 with Fig. 9, it is likely that 383 the amorphous phases are responsible for the compressive strength. The incorporation of PG gave 384 rise to thenardite formation, which increased the GGBFS dissolution and consequently the merwinite 385 content after 28 days was lower compared to the reference samples. Secondary gypsum is present 386 for SS1, SH2 and SH3 upon the dissolution of early ettringite, however does not impede a negative 387 influence on the compressive strength after 28 days. Nonetheless, compressive strength testing and 388 phase development after prolonged curing times should be the subject of further investigation.

389

## 390 4. Conclusions

391 In this study, the effect of PG incorporation in alkali-activated GGBFS on the reactivity, strength 392 development and final matrix properties was investigated, as a function of alkali activator. The 393 results were compared with alkali-activated reference samples based on GGBFS. PG completely 394 dissolved and took part in solid reaction product formation; a hydraulic binder was created. Upon PG 395 incorporation, portlandite and ettringite were formed immediately after dissolution of precursors. 396 The hardened binder consisted mainly of amorphous hydration products, intermixed with thenardite 397 and minor amounts of secondary gypsum and undissolved merwinite. The formation of thenardite 398 was enviable from the microstructural development point of view, giving rise to accelerated GGBFS 399 dissolution. However, leftovers of thenardite may result in high leaching rates of sodium and 400 sulphate when in contact with water. PG incorporation resulted in faster initial setting (i.e. a reduction of 0.1 h to 1.2 h) and delayed final setting time (i.e. an increase of 0.1 h to 2 h), compared 401 402 with the reference samples. The incorporation of PG enhanced the compressive strength with 55.3% 403 to 89.0% at 28 days when sodium hydroxide was used as alkali activator, likely due to an increase in 404 the amorphous content. Both the compressive strength and amorphous content were reduced when 405 incorporating PG in sodium silicate activated samples. Regions of secondary gypsum and thenardite 406 were clearly detectable in micro-chemical analysis. An amorphous C-A-S-H type gel was the prevalent 407 reaction product and the incorporation of PG gave rise to a decrease of Al/Si and Ca/Si ratios in the 408 C-A-S-H gels and a higher polymerized network.

409

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