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# **1** Pore solution composition of alkali-activated slag/fly ash pastes

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Abstract: The pore solutions of a series of hardened alkali-activated slag/fly ash pastes were 8 extracted by the steel-die method, and analyzed using ICP-OES analysis technique. According to 9 the saturation index from thermodynamic calculations, the pore solutions of alkali-activated slag 10 pastes kept oversaturated with respect to solid reaction products with time. In the pore solutions 11 of alkali-activated fly ash pastes, an increase of temperature (from 40 °C to 60 °C) led to 12 decreases of the concentrations of Si, Al, Ca, Na, OH, K, Fe and Mg, while the soluble silicate in 13 the alkaline activator resulted in increases of the concentrations of these elements. Compared to 14 the alkali-activated slag paste with the same alkaline activator, 50% replacement of slag by fly 15 ash did not result in a substantial change of the pore solution composition. Based on the 16 experimental results, conceptual models were proposed to describe the elemental concentrations 17 in the pore solutions. 18

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20 Keywords: alkali activation; slag/fly ash; pore solution; saturation index; conceptual models

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## 25 **1 Introduction**

26

Alkali-activated materials (AAMs) are a class of materials obtained by the chemical reaction of a 27 powdered aluminosilicate precursor (most commonly blast furnace slag and/or fly ash) with an 28 29 alkaline activator (usually a solution of sodium hydroxide and/or sodium silicate) at ambient or elevated temperature [1, 2]. Due to the ability to utilize high-volume of industrial by-products as 30 the powdered aluminosilicate precursors, AAMs can save a large amount of fossil fuel-derived 31 energy. As such, 80% or greater reduction of CO<sub>2</sub> emission can be achieved by AAMs when 32 compared to ordinary Portland cement (OPC) based materials [3, 4]. Besides, AAMs release 33 lower amount of heat [5], and provide comparable or even superior mechanical properties and 34 durability performance [3, 6-9]. Therefore, AAMs show very promising potential for utilization 35 to embrace a sustainable future global construction materials industry [2]. 36

37

38 Pore solution is an essential component of AAMs. It is in this medium where the powdered 39 aluminosilicate precursors dissolve and subsequently a set of reactions take place to produce solid 40 reaction products. The role of pore solution can be viewed in the following three aspects.

41

*Dissolution kinetics.* The high concentration of OH<sup>-</sup> ions in the pore solution allow
 breaking of bonds such as Si-O and Al-O on the surface of aluminosilicate precursors [10].
 A higher alkalinity of pore solution will accelerate the dissolution kinetics [11-15].
 Therefore, from the kinetics point of view, pore solution composition determines the
 dissolution kinetics of aluminosilicate precursors. The dissolution kinetics determines the
 degree of reaction and thus controls the reaction kinetics and the amount of solid reaction
 products produced with time.

49

Thermodynamics. Thermodynamics is of great significance to the understanding of 50 chemical reactions [16]. The reactions of AAMs, same as any other chemical system, 51 52 follow the law of thermodynamics. When aluminosilicate precursors are brought into contact with an alkaline activator, their constituents start to dissolve and subsequently 53 various solid reaction products start to be formed if the solution is saturated or 54 oversaturated with respective the solids. At the given temperature and pressure, the pore 55 solution composition controls the ongoing reaction process and determines what reaction 56 products are to be formed [17, 18]. 57

58

Durability. The dominant phase assemblage in AAMs is an alkali calcium-aluminosilicate 59 hydrate (C-(N-)A-S-H) type gel or a three-dimensional hydrous alkali-aluminosilicate (N-60 A-S-H) type gel, depending on the Ca content in the system [1]. From the durability point 61 of view, the high alkalinity of pore solution ensures the stability of calcium-62 aluminosilicate hydrates [19] and alkali-aluminosilicate gels [20] in AAMs. If AAMs are 63 embedded with reinforcing steel, the pore solution plays a more important role by forming 64 65 a passive film on the surface of steel rebars and, consequently, avoiding corrosion of steel rebar [21, 22]. 66

67

Those three aspects illustrate the significance of pore solution in AAMs. Furthermore, the chemical properties (chemical composition, and amorphous content etc.) and physical properties (surface morphology, and particle size etc.) of aluminosilicate precursors vary significantly from different sources [1, 5, 23]. For this reason, the alkali activation of aluminosilicate precursors greatly relies on the pore solution. However, studies regarding the pore solution composition in

AAMs are few for alkali-activated slag paste [12, 24-26], and even rarely reported for alkaliactivated fly ash paste and alkali-activated slag blended with fly ash paste.

75

Song and Jennings studied the pore solution chemistry of alkali-activated slag and found that 76 77 higher alkalinity of pore solution led to higher concentrations of Si and Al and lower concentrations of Ca and Mg [24]. In another study regarding the relationship between the 78 composition of pore solution and the composition and structure of the main reaction products in 79 alkali-activated slag paste, the researchers found that the nature of the alkaline activator 80 influenced the structure and composition of the produced calcium silicate hydrate [25]. In 81 comparison with the pore solution of OPC-based materials, the pore solution of alkali-activated 82 slag paste showed ten times lower concentration of Ca and tens to hundreds of times higher 83 concentrations of Si and Na [12]. It was reported that the initiation of pitting at the steel-cement 84 concrete interface started only after the concentration ratio of Cl<sup>-</sup> relative to OH<sup>-</sup> exceeded a 85 particular value [22]. This suggests a great dependence of passivation of reinforcing steel on the 86 alkalinity of the pore solution. Because of little soluble Ca available in the pore solution of alkali-87 activated slag to play a buffering role similar to Ca(OH)<sub>2</sub> in OPC-based materials, the alkalinity 88 of pore solution of alkali-activated slag is mainly maintained by the content of alkali [26]. 89 Therefore, it is essential to provide sufficient amount of alkalis to ensure protection of reinforcing 90 steel from corrosion. 91

92

93 In this study, the pore solutions of alkali-activated slag, alkali-activated fly ash and alkali-94 activated slag blended with fly ash pastes were studied by means of the inductively coupled 95 plasma-optical emission spectroscopy (ICP-OES) analysis technique. The influences of alkaline 96 activator and temperature on the pore solution composition were investigated. The measured

97 composition of the pore solution and its changes with time can be used to predict the (potentially) 98 solid reaction products. Thus, the pore solutions of alkali-activated slag pastes were 99 thermodynamically analyzed in terms of effective saturation index. In the meantime, solid phase 100 analysis was conducted through the X-ray diffraction (XRD) and Fourier transform infrared 101 spectroscopy (FTIR). Based on the experimental results, conceptual models were proposed to 102 describe the elemental concentrations in the pore solutions. The obtained results will contribute to 103 new insights regarding the chemistry of pore solution and thermodynamic modelling in AAMs.

104

#### 105 2 Materials and methods

106

107 2.1 Materials and mixtures

108

In this study, ground granulated blast furnace slag and fly ash were used as the aluminosilicate 109 precursors to prepare alkali-activated slag/fly ash pastes. The chemical compositions of blast 110 furnace slag and fly ash were determined by X-ray fluorescence spectrometry (XRF), and are 111 112 listed in Table 1. The X-ray diffraction patterns of slag and fly ash are plotted in Figure 1. Quartz, mullite and hematite were the three crystalline phases identified in fly ash. In order to determine 113 the amorphous content of fly ash, corundum ( $\alpha$ -Al<sub>2</sub>O<sub>3</sub>) was added as an internal standard to 114 replace 20% of fly ash. Through the Rietveld analysis method [27], the amorphous content of fly 115 ash was determined as 71%. By subtracting SiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub> in the crystalline phases from the 116 total SiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub>, the reactive SiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub> were determined as 37% and 15% respectively. 117 These results agree with the measurements by a previous colleague using the chemical 118 dissolution treatment (we used a similar fly ash from the same manufacture plant in Netherlands) 119 [28]. For slag, no crystalline phase was identified from the X-ray diffraction pattern. This is 120

121 consistent with the fact that the ground granulated blast furnace slag usually contains more than122 95% vitreous phase [29].

123

**124** Table 1 Chemical compositions of blast furnace slag and fly ash (by weight, %)

125

Figure 1 X-ray diffraction patterns of slag and fly ash. In the graph, Q, M and H refer to quartz, mullite and hematiterespectively.

128

Sodium hydroxide (analytical grade, >98%) and water glass (8.25 wt.% Na<sub>2</sub>O, 27.5 wt.% SiO<sub>2</sub> 129 and 64.25 wt.% H<sub>2</sub>O) were used to prepare two types of alkaline activators: sodium hydroxide 130 activator and sodium silicate activator. As shown in Table 2, a series of alkaline activators with 131 different contents of NaO<sub>2</sub> and SiO<sub>2</sub> were used to prepare alkali-activated slag pastes, alkali-132 activated fly ash pastes and alkali-activated slag blended with fly ash pastes. In the notations for 133 the samples, N and S indicate weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to the precursor, 134 T indicates temperature in Celsius and B indicates blend of slag and fly ash. The Na<sub>2</sub>O in alkaline 135 activator consists of the Na<sub>2</sub>O from sodium hydroxide and the Na<sub>2</sub>O from water glass. For alkali-136 activated slag pastes and alkali-activated slag blended with fly ash pastes, water to precursor ratio 137 was 0.4. For alkali-activated fly ash pastes, water to precursor ratio was 0.35. Here, the water in 138 139 water glass was also included in the calculation of the water to precursor ratio of the pastes. According to the type of alkaline activator, the samples were categorized into two systems: 140 sodium hydroxide activated system (with no soluble silicate in the alkaline activator) and sodium 141 142 silicate activated system (with soluble silicate in the alkaline activator).

143

144 Table 2 Mixture compositions and curing temperature

According to the materials characterization described previously, fly ash contains glassy 146 (amorphous) and crystalline constituents in nature, e.g. guartz, mullite and hematite. Compared to 147 slag, fly ash has much lower reactivity. Therefore, elevated temperature curing is usually applied 148 for the alkali activation of fly ash in order to obtain a relatively high strength at early ages [28, 149 150 30]. The elevated temperature leads to an acceleration of the dissolution of fly ash and the polymerization process of alkali-aluminosilicate type gel [1, 31]. In this study, two elevated 151 temperatures (40 °C and 60 °C) were used to investigate the influence of temperature on the pore 152 solution composition. For all samples, sealed curing was applied until testing. 153

154

The pH and concentrations of Na, Si, and OH<sup>-</sup> in the alkaline activators are listed in Table 3. 155 These results were obtained through the thermodynamic calculations using a thermodynamic 156 software. This software is introduced in the section 2.3. In sodium hydroxide solution, aqueous 157 sodium hydroxide (NaOH(aq)) normally exists. For this reason, the measured concentration of 158 OH<sup>-</sup> through titration against hydrochloride acid includes the hydroxide from NaOH(aq). The 159 details of titration against hydroxide acid are given in section 2.2. As derived by thermodynamic 160 161 calculations, two values of the concentration of OH<sup>-</sup>, e.g. including and excluding hydroxide from NaOH(aq), are provided in Table 3. A big difference can be seen between those two values for 162 each alkaline activator, and the concentration of OH<sup>-</sup> including hydroxide from NaOH(aq) is 163 closer to that in the pore solution of alkali-activated slag/fly ash pastes, as obtained by titration. 164

165

166 Table 3 The pH and concentrations of Na, Si, and OH<sup>-</sup> in the alkaline activators

167

168 2.2 Pore solution analysis via ICP-OES

The pore solutions of hardened samples were extracted using the steel-die method as described by Barneyback and Diamond [32]. Pressures of up to 300 MPa were used to extract the pore fluid from the samples at different curing ages. After extraction, the pore solution was immediately filtered using Whatman 41 filter paper. One part of the filtered solution was diluted using nitric acid (0.2 vol.%). The diluted solutions were analyzed through a PerkinElmer Optima 5300DV ICP-OES spectrometer, by which the concentrations of Si, Al, Ca, Na, K, Fe and Mg were determined.

177

The concentration of OH<sup>-</sup> was measured by titration against hydrochloride acid (0.1 mol/L). 178 Phenolphthalein was used as the indicator. First, a volume  $(V_1, ml)$  of pore solution was 179 accurately measured using a pipette and injected into a beaker. Then a drop of phenolphthalein 180 was added into the beaker, and the solution turned red. Afterwards, hydrochloride acid was 181 slowly added into the beaker through a burette until the solution turned colorless. While adding 182 hydrochloride acid, the beaker was swirled with hand. After the titration, the amount of 183 hydrochloride acid added into the beaker was measured as  $V_2$  (ml). Then the concentration of 184 OH<sup>-</sup> ([OH<sup>-</sup>], mol/L) could be determined as: 185

186

187  $[OH^-] = 0.1 * V_2 / V_1$  (1)

188

In the titration test, two replicates of each pore solution were titrated, and the average [OH<sup>-</sup>] was
presented as the final result.

191

192 For the measurement of S, another part of the filtered solution was diluted using sodium193 hydroxide solution (0.1 mol/L) instead of nitric acid, and subsequently subject to ICP-OES

194	analysis. This is because S mostly exists as sulfide (HS <sup>-</sup> ) in the pore solution [12, 33]. The
195	dilution with nitric acid will acidify $HS^-$ and lead to formation of $H_2S$ as described by Equation
196	(2). On one hand, $H_2S$ evaporates and results in a characteristic foul odor of rotten eggs, as
197	experienced during the dilution process using nitric acid. On the other hand, $H_2S$ may be oxidized
198	and lead to white precipitation of sulfur as described by Equation (3), and shown in Figure 2.
199	
200	$\mathrm{HS}^{-} + \mathrm{H}^{+} \to \mathrm{H}_{2}\mathrm{S}. \tag{2}$
201	
202	$2H_2S + O_2 \rightarrow S_2 \downarrow + 2H_2O \tag{3}$
203	
204	Figure 2 shows the white precipitation under the scanning electron microscope (SEM) and its
205	element composition using energy dispersive spectroscopy. This finding confirmed the fact that
206	acid dilution resulted in white precipitation of sulfur. For this reason, the measured
207	concentrations of S using nitric acid dilution for alkali-activated slag paste were not correct and
208	thus not presented in this study.
209	
210	Figure 2 White precipitation and its element composition in diluted pore solution using nitric acid.
211	
212	2.3 Thermodynamic analysis of pore solution
213	
214	Thermodynamics is essential to the understanding of chemical reactions. Whether a solid can
215	
	potentially form or not can be predicted through the saturation index (SI) [17].
216	potentially form or not can be predicted through the saturation index (SI) [17].

218

where IAP and  $K_{S0}$  are the ion activity product and equilibrium solubility product of a solid respectively. A positive SI indicates oversaturation and thus the possibility that this solid phase can precipitate. If SI is negative, it means under-saturation and implies that this solid is unstable in equilibrium within such a solution. A SI of zero indicates equilibrium between solution and solid. However, SI can be misleading when comparing phases which dissociate into a different number of ions (N) [17]. For this reason, effective saturation index (ESI) is applied in this study for thermodynamic analysis of pore solution.

226

 $227 \quad ESI = SI/N \tag{5}$ 

228

Before performing the thermodynamic analysis of the pore solution, it is crucial to obtain the 229 thermodynamic database. In AAMs, the thermodynamic database is only available for alkali-230 activated slag. For alkali-activated fly ash, the thermodynamic data have not been established yet. 231 For this reason, the thermodynamic analysis was only performed for the pore solutions of the 232 alkali-activated slag pastes. The Gibbs energy minimization software GEM-Selektor v.3 233 (http://gems.web.psi.ch/) [34, 35] and the thermodynamic database developed in [36, 37] for 234 alkali-activated slag were used. The activities of ions were calculated using the measured 235 elemental concentrations as the input. 236

237

238 2.4 Measurement of heat evolution rate

239

The heat evolution rate was measured using an isothermal calorimeter by following standard
ASTM C1679 [38]. For the measurements of heat evolution at the elevated temperatures (40 °C

and 60 °C), internal mixing was used to avoid any temperature difference. Fly ash and alkaline
activator were conditioned in the calorimeter at the measurement temperature. When the thermal
equilibrium was reached, the alkaline activator was injected into the glass ampoule to be mixed
with aluminosilicate precursor powder internally. Subsequently, the heat release rate was
recorded.

247

248 2.5 X-ray diffraction (XRD) and Fourier transform infrared spectroscopy (FTIR)

249

250 XRD analysis was performed on the powdered samples using a Philips X'Pert diffractometer 251 applying CuK<sub> $\alpha$ </sub> radiation ( $\lambda = 1.54$  Å). The samples were scanned between 5° and 70° 2-theta, 252 with a step size of 0.02° 2-theta and a dwell time of 5 s per step. FTIR was performed using a 253 Spectrum TM 100 Optical ATR-FTIR spectrometer over the wavelength range of 600 to 4000 254 cm<sup>-1</sup> with a resolution of 4 cm<sup>-1</sup>.

255

256 3 Results and discussion

257

- 258 3.1 Pore solution composition of alkali-activated slag paste
- 259

Figure 3 presents the concentrations of Si, Al, Ca, OH<sup>-</sup>, K and Na (hereafter denoted as [Si], [Al], [Ca], [OH<sup>-</sup>], [K] and [Na] respectively) in the pore solution of alkali-activated slag paste as a function of time up to 28 days. The pore solution data are listed in the Appendix. The pore solution composition was dominated by Na and OH<sup>-</sup>, whereas much lower concentrations of Si, Al, K and Ca were observed. The most significant changes in the compositions of pore solutions in alkali-activated slag pastes took place during the first hours up to 3 days. This can be seen

from the changes of elemental concentrations with time, particularly the decreases of the [Si], [Na] and  $[OH^-]$ . The concentrations of Si, Al and Ca were in the order: [Si] > [Al] > [Ca]. The developments of the elemental concentrations with time will be further discussed in the section 3.5.

270

Figure 3 Elemental concentrations in the pore solutions of alkali-activated slag pastes. In the graphs, N and S
indicate weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The water to slag ratio was 0.4 and the curing
temperature was 20 °C.

274

There was a strong decrease of [Si] during the initial period up to 3 days, and then the decrease of 275 [Si] continued gradually with time for all samples. For sodium hydroxide activated slag system, 276 277 the [Si] increased with the increase of Na<sub>2</sub>O content. For sodium silicate activated system, the 278 [Si] increased with the increase of Na<sub>2</sub>O content after 1 day. The [Si] in the pore solution of 279 sodium silicate activated slag paste was much higher than that in the sodium hydroxide activated 280 slag paste with the same content of Na<sub>2</sub>O. The higher [Si] for sodium silicate activated slag paste was a result of the soluble Si in the sodium silicate activator. For all the alkali-activated slag 281 samples, the [A1] decreased slightly with time. The increase of Na<sub>2</sub>O content in alkaline activator 282 led to an obvious increase of the [Al]. 283

284

The  $[OH^-]$  decreased over time and increased with the increase of Na<sub>2</sub>O content. Comparing the sodium silicate activated sample to the sodium hydroxide activated sample with the same content of Na<sub>2</sub>O, it can be seen that the added SiO<sub>2</sub> from alkaline activator resulted in a decrease of  $[OH^-]$ . The  $[OH^-]$  indicates a pH in the range of 13.77~14.67 up to 28 days in the pore solution. This shows a higher alkalinity than that in the pore solution (pH>12) buffered by Ca(OH)<sub>2</sub> in OPC-



based materials, and much higher than the pH of 9.5 below which corrosion commences [39].

291 The [Ca] remained at a very low level (0.2~2 mmol/L) and did not show much change over time.

292

An apparent decrease of [Na] was observed during the first 3 days in Figure 3(E). And then the [Na] decreased slightly with progressive reaction of slag. When the Na<sub>2</sub>O content increased from 4% to 8%, the [Na] increased for both the sodium hydroxide activated slag system and the sodium silicate activated slag system. On the contrary to the [Na], the [K] increased with time. This was because K was sourced from slag and increased with the dissolution of slag, while Na was mainly sourced from the alkaline activator and decreased as it was taken up into the calcium aluminosilicate hydrates [33, 40, 41].

300

In the pore solution of alkali-activated slag, the concentrations of Mg and Fe were not presented 301 since they were below the detection limit of ICP-OES. The detection limit of Mg and Fe are 302 0.0042 mmol/L and 0.0009 mmol/L respectively. The concentration of Mg below the detection 303 limit was attributed to the formation of a hydrotalcite-like phase in alkali-activated slag as seen 304 305 from the XRD patterns (see Figure 8). The formation of hydrotalcite led Mg within the pore solution into hydrotalcite, and thus resulted into low concentration of Mg. The concentration of 306 Fe below the detection limit was because of the low content of  $Fe_2O_3$  in slag (0.46%). On the 307 contrary, the concentrations of Mg and Fe were obtained for the alkali-activated fly ash pastes, 308 309 which will be presented in section 3.3. This was because of no Mg containing phase detected in alkali-activated fly ash (see Figure 11) and much higher content of  $Fe_2O_3$  in fly ash (6.6%). 310

311

The plots of the concentrations of Na, Al and Ca as a function of the concentration of OH<sup>-</sup> are shown in Figure 4(A). In pore solutions, the positive charge of Na is mostly balanced by the

negative hydroxide ion. Higher [Na] leads to higher [OH]. For this reason, the concentration of 314 315 Na increased linearly with the increase of the concentration of OH<sup>-</sup> and this relationship was 316 independent of the type of alkaline activator. In pore solutions, the concentration of Ca remained at a very low level and slightly decreased as the concentration of OH<sup>-</sup> increased. This could be 317 318 attributed to the common ion effect [42, 43]. The pore solutions had high concentrations of Si, OH<sup>-</sup> and Na<sup>+</sup>. Those ions might combine with Ca to form solid reaction products (see Table 4) 319 and thus lowered the Ca concentration in the pore solution. This point could be further supported 320 321 by thermodynamic analysis of pore solution (see section 3.2). The ESI with respect to C-(N-)A-S-H and secondary reaction products were mostly larger than zero, which indicates oversaturation 322 and likely precipitation of these solid phases in pore solution. Therefore, Ca stayed on a low 323 concentration and the increase of [OH<sup>-</sup>] led to a decrease of the [Ca] in the pore solution. 324

325

The [A1] increased with the increase of [OH] in the pore solution for both the sodium hydroxide 326 activated slag system and the sodium silicate activated slag system, which is consistent with the 327 finding in the literature [24]. However, this relationship only exists for the samples using the 328 329 same type of alkaline activator. Using the alkaline activator with the same content of Na<sub>2</sub>O, the sodium hydroxide activated slag paste (e.g. N4S0) had higher [OH<sup>-</sup>] in the pore solution than that 330 of the sodium silicate activated slag paste (e.g. N4S5.4). But the pore solution of sodium 331 hydroxide activated slag paste (e.g. N4S0) had a lower [A1] than that of sodium silicate activated 332 slag paste (e.g. N4S5.4). This was because of the different calcium aluminosilicate hydrates 333 formed as a consequence of the different types of alkaline activators used in the alkali activation 334 of slag. The calcium aluminosilicate hydrate formed in the sodium hydroxide activated slag paste 335 had a higher structural order than that formed in the sodium silicate activated slag paste [25]. This 336 is also found in this work by XRD analysis (see Figure 8). For sodium hydroxide activated 337

samples (for instance N4S0), the peak that corresponded to C-(N-)A-S-H, was sharper and had a 338 higher intensity than that for sodium silicate activated slag samples (for instance N4S5.4). 339 Furthermore, C-S-H(I) was detected in sodium hydroxide activated slag samples while it was not 340 in sodium silicate activated slag samples. As such, the calcium aluminosilicate hydrate formed in 341 342 sodium hydroxide activated slag had higher structural order and degree of crosslinking. The higher structural order and degree of crosslinking of calcium aluminosilicate hydrate stabilized its 343 structure, which could lead to a lower solubility. The lower solubility of calcium aluminosilicate 344 hydrate resulted in lower solubility of Al and thus lower [Al] in the pore solution of sodium 345 hydroxide activated slag paste. 346

347

Figure 4 Concentrations of one element as a function of the concentration of another element in pore solution. (A):
[Na], [Al] and [Ca] versus [OH<sup>-</sup>]. (B): [Si] versus [OH<sup>-</sup>]. (C) [Si] versus [Ca]. In the graphs, N and S indicate the
weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The curing temperature was 20 °C. The water to slag ratio
was 0.4.

352

In alkaline solutions, the solubility of Si increases with the increase of the alkalinity of solution 353 [11]. This is also observed in the pore solution of sodium hydroxide activated slag paste [24]. In 354 this study, the relationship between the alkalinity of pore solution and the solubility of Si was 355 more comprehensively studied for sodium hydroxide activated slag pastes and sodium silicate 356 activated slag pastes. Figure 4(B) shows the plots between the concentration of Si and the 357 358 concentration of OH<sup>-</sup> in the pore solution. The [Si] increased with the increase of [OH<sup>-</sup>] in the pore solution for each alkali-activated slag paste. The plots suggest the existence of two separate 359 lines as shown in Figure 4(B). The line with a higher [Si] is associated with the sodium silicate 360 activated slag pastes, while the line with a lower [Si] corresponds to the sodium hydroxide 361

activated slag pastes. From the sodium hydroxide activated slag system to the sodium silicate activated slag system, the concentration of Si moves up and backward as indicated by the arrows. This is because of the addition of SiO<sub>2</sub> from the sodium silicate activator. On one hand, the addition of SiO<sub>2</sub> resulted in higher [Si] in the pore solution and thus an up translation of the plot. On the other hand, the addition of SiO<sub>2</sub> resulted in lower alkalinity of pore solution, e.g. lower [OH<sup>-</sup>], and thus a backward translation of the plot.

368

Figure 4(C) plots the concentration of Si as a function of the concentration of Ca. As marked with a dashed black line in the graph, a decreasing trend was generally observed between the concentrations of Si and Ca, which is in line with the literature [24, 44]. The fact that the concentration of Si decreased with the increasing concentration of Ca was attributed to the common ion effect. Excess of Ca within pore solution could combine Si into solid reaction products and thus result in a lower concentration of Si.

375

376 3.2 Thermodynamic analysis of the pore solution of alkali-activated slag paste

377

As it is directly encoded in GEM-Selektor, the extended Debye-Huckel equation is used to 378 calculate the ion activity coefficients [34, 35]. The extended Debye-Huckel equation is accurate 379 at moderate ionic strengths (up to ~1-2 molal) [45]. Figure 5 presents the ionic strengths in the 380 pore solution of alkali-activated slag paste. The ionic strengths in the pore solutions of N4S0, 381 N4S5.4 and N6S5.4 were within 1-2 molal. In the pore solutions of N6S0, N8S0 and N8S5.4, the 382 ionic strengths were larger than 2 molal, beyond the valid range (~1-2 molal). However, use of an 383 improved aqueous phases model, such as the Pitzer model [46], is limited in the current version 384 of GEM-Selektor. This is because the description of aqueous silicate species in the database of 385

GEM-Selektor does not extend beyond dimeric silicate and aluminosilicate units [36]. On the 386 387 other hand, GEM-Selektor has been commonly used in the thermodynamic modelling of the hydration of alkali-activated slag (with ionic strengths  $\sim 1-3$  molal in the pore solution [12]), and 388 the modelling results agreed well with the experimental data [37, 47, 48]. As such, slight excess 389 390 of ionic strengths (> 2 molal) would not impose a serious impact on the calculation results. Since the measured elemental concentrations, including the concentration of OH, were used as input to 391 calculate the activities of aqueous ions with GEM-Selektor, the calculated pH agreed with the 392 measured one. 393

394

Figure 5 The ionic strengths in the pore solutions of alkali-activated slag pastes, calculated by GEM-Selektor. In the
graphs, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The curing temperature was
20 °C. The water to slag ratio was 0.4.

398

Myers et al. used a set of eight CNASH\_ss end-members to describe the calcium aluminosilicate 399 400 hydrates in alkali-activated slag [36]. These eight end-members and their corresponding dissociation reactions and equilibrium solubility products are presented in Table 4. Among the 401 eight end-members for the calcium aluminosilicate hydrates, 5CA and INFCA are two C-A-S-H 402 gel end-members, 5CNA and INFCNA are two C-N-A-S-H gel end-members, INFCN is a C-N-403 S-H gel end-member, and T2C\*, T5C\* and TobH\* are three C-S-H gel end-members. The 404 equilibrium solubility products of these eight end-members were used to calculate the effective 405 saturation indexes as described in section 2.3. 406

407

Table 4 Chemical reactions and equilibrium solubility products at 25°C and 1 bar for eight end-members and
 crystalline reaction products in alkali-activated slag

410

Figure 6 presents the effective saturation indexes with respect to the eight end-members. For the 411 sodium hydroxide activated slag pastes (the left column in Figure 6), the pore solutions were 412 oversaturated with respect to 5CA, 5CNA, INFCNA, INFCN, T2C\* and T5C. The effective 413 414 saturation indexes with respect to those end-member gels decreased with time during the initial period (up to 3 days). This indicates the continuous formation of the calcium aluminosilicate 415 hydrates. The formation of the calcium aluminosilicate hydrates reduced the degree of 416 417 oversaturation and thus led to decreases of the effective saturation indexes. After the initial period, the reaction of slag slowed down and stepped into a steady reaction stage controlled by 418 diffusion processes [49]. In this reaction stage, there might exist a dynamic equilibrium between 419 the dissolution of slag and the formation of the calcium aluminosilicate hydrates. For this reason, 420 the effective saturation indexes with respect to 5CA, 5CNA, INFCNA, INFCN, T2C\* and T5C 421 showed no substantial change with time. Although the increase of Na<sub>2</sub>O content resulted in 422 higher alkalinity of the pore solution, it did not lead to significant influence on the effective 423 saturation indexes with respect to the eight end-members. 424

425

For sodium silicate activated slag pastes (the right column in Figure 6), similar results were found. Compared to those for the sodium hydroxide activated slag pastes, the effective saturation indexes with respect to the eight end-members were slightly higher for the sodium silicate activated slag pastes. The slightly higher effective saturation indexes were attributed to the larger activity coefficients of  $SiO_3^{2-}$  and  $Ca^{2+}$  (two valence electrons) than those of Na<sup>+</sup> and OH<sup>-</sup> (one valence electron). It can be seen in Figure 6 that the pore solution of sodium silicate activated slag paste had higher concentrations of Si, Al and Ca than that of sodium hydroxide activated slag paste. As such, the pore solution of sodium silicate activated slag paste had higher effectivesaturation indexes with respect to the eight end-members.

435

**436** Figure 6 Effective saturation indexes with respect to the eight end-members as functions of time. In the graphs, N 437 and S indicate the weight percentage of  $Na_2O$  and  $SiO_2$  with respect to slag. The curing temperature was 20 °C. The 438 water to slag ratio was 0.4. A ESI of 0 indicates equilibrium between solution and solid; a positive ESI indicates 439 oversaturation and a negative ESI indicates undersaturation.

440

In addition to the calcium aluminosilicate hydrates, some crystalline reaction products are also 441 442 identified in alkali-activated slag, such as hydrotalcite [50], tetracalcium aluminate hydrate (C<sub>4</sub>AH<sub>13</sub>) [50], katoite (C<sub>3</sub>AH<sub>6</sub>) [37] and stratlingite (C<sub>2</sub>ASH<sub>8</sub>) [51]. The dissociation reactions 443 and equilibrium solubility products of C<sub>2</sub>ASH<sub>8</sub>, C<sub>3</sub>AH<sub>6</sub>, C<sub>4</sub>AH<sub>13</sub> and portlandite (Ca(OH)<sub>2</sub>) are 444 listed in Table 4. These products were also captured according to the effective saturation index 445 from thermodynamic calculations. Figure 7 plots the effective saturation indexes with respect to 446 C<sub>2</sub>ASH<sub>8</sub>, C<sub>3</sub>AH<sub>6</sub>, C<sub>4</sub>AH<sub>13</sub> and Ca(OH)<sub>2</sub> for the sodium hydroxide activated slag pastes (left 447 column) and the sodium silicate activated slag pastes (right column). Since the concentration of 448 Mg was not measured, the effective saturation index with respect to hydrotalcite was not 449 calculated. The effective saturation indexes with respect to C<sub>2</sub>ASH<sub>8</sub>, C<sub>3</sub>AH<sub>6</sub>, C<sub>4</sub>AH<sub>13</sub> and 450 Ca(OH)<sub>2</sub> increased rapidly during the initial period. This is in the contrast to the effective 451 saturation indexes with respect to 5CA, 5CNA, INFCNA, INFCN, T2C\* and T5C. It indicates 452 453 that the calcium aluminosilicate hydrate is thermodynamically favored to form at early age, while the crystalline reaction products are thermodynamically favored to form at a later age. After the 454 initial period, the effective saturation indexes with respect to these four crystalline phases showed 455 no substantial change with time. The effective saturation indexes with respect to these four 456

457 crystalline phases were in the order:  $ESI(C_2ASH_8) > ESI(C_3AH_6) > ESI(Ca(OH)_2) \ge$ 458  $ESI(C_4AH_{13})$  for the sodium hydroxide activated slag pastes, and  $ESI(C_2ASH_8) > ESI(C_3AH_6) >$ 459  $ESI(C_4AH_{13}) \ge ESI(Ca(OH)_2)$  for the sodium silicate activated slag pastes.

460

Figure 7 Effective saturation indexes with respect to C<sub>2</sub>ASH<sub>8</sub>, C<sub>3</sub>AH<sub>6</sub>, C<sub>4</sub>AH<sub>13</sub> and Ca(OH)<sub>2</sub> as functions of time. In
the graphs, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The curing temperature
was 20 °C. The water to slag ratio was 0.4. A ESI of 0 indicates equilibrium between solution and solid; a positive
ESI indicates oversaturation and a negative ESI indicates undersaturation.

465

466 In order to confirm the thermodynamic analysis results, solid phase analysis was conducted for alkali-activated slag pastes using XRD. Figure 8 presents the XRD patterns of N4SO, N8SO and 467 N4S5.4 at 28 days. It can be seen that hydrotalcite and katoite (C<sub>3</sub>AH<sub>6</sub>) were identified in all 468 samples, which is in line with previous studies [50, 52, 53]. As the Na<sub>2</sub>O content increased from 469 4% to 8%, portlandite was detected. The diffusive peak at  $2\theta = 29.07^{\circ}$  was dominant in all alkali-470 activated slag samples. This diffusive peak is associated with the calcium aluminosilicate 471 hydrates (C-(N-)A-S-H) [52, 54]. According to the previous studies [50, 52, 54], the peaks at 472 approximately  $2\theta = 7.0^{\circ}$ ,  $32.0^{\circ}$  and  $49.8^{\circ}$  are attributed to the poorly crystalline C-S-H(I). The 473 poorly crystalline C-S-H(I) is considered more ordered than the C-S-H in ordinary Portland 474 cement paste at ambient temperature [55] and has been observed in alkali-activated slag [56]. It is 475 noted that C-S-H(I) was only detected in sodium hydroxide activated slag samples. The XRD 476 results validated the thermodynamic analysis using effective saturation index. The solid reaction 477 products with positive effective saturation indexes were likely to precipitate, such as calcium 478 aluminosilicate hydrate, katoite and portlandite. 479

Figure 8 X-ray diffraction patterns of N4S0, N8S0 and N4S5.4 at 28 days. In the graphs, N and S indicate the weight
percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The curing temperature was 20 °C. The water to slag ratio was 0.4.

484 3.3 Pore solution composition of alkali-activated fly ash paste

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Figure 9 presents the measured concentrations of Si, Al, OH<sup>-</sup>, Ca, Na, K, Fe, Mg and S in the 486 pore solution of alkali-activated fly ash pastes as a function of time up to 28 days (672 hours). 487 The pore solution data are listed in the Appendix. The pore solution composition was dominated 488 by Na, OH<sup>-</sup> and Si, whereas much lower concentrations of Al, K, Ca, Fe and Mg were observed. 489 The most significant changes in the compositions of pore solutions in alkali-activated fly ash 490 pastes took place during the first week. This can be seen from the changes of elemental 491 concentrations with time, particularly the decreases of the [Si], [Na] and [OH<sup>-</sup>]. The 492 concentrations of Si, Al and Ca were in the order: [Si] > [Al] > [Ca]. The developments of the 493 494 elemental concentrations with time will be further discussed in the section 3.5. The increase of curing temperature from 40 °C to 60 °C led to decreases of the concentrations of Si, Al, OH, Ca, 495 Na, K, Fe and Mg. The influence of temperature on the pore solution composition of alkali-496 497 activated fly ash paste will be discussed in the section 3.6.

498

**Figure 9** Elemental concentrations in the pore solutions of alkali-activated fly ash pastes where N and S indicate the weight percentage of  $Na_2O$  and  $SiO_2$  with respect to fly ash, and T indicates the curing temperature in Celsius. The water to fly ash ratio was 0.35.

502

It can been seen from Figure 9(A) that the [Si] decreased with time for all alkali-activated fly ash
pastes. As the Na<sub>2</sub>O content increased from 6.2% to 9.3%, the [Si] increased for both curing

temperatures. The [Si] was much higher in the sodium silicate activated fly ash paste than that in the sodium hydroxide activated fly ash paste with the same content of  $Na_2O$ . This was attributed to the soluble silica added from the alkaline activator prepared using sodium silicate solution.

As shown in Figure 9(B), the [Al] decreased with time for all alkali-activated fly ash samples. For the sodium hydroxide activated fly ash pastes, the [Al] increased slightly as the Na<sub>2</sub>O content increased from 6.2% to 9.3%. When the sodium silicate was added in the alkaline activator, the [Al] increased significantly in comparison with the sodium hydroxide activated fly ash with the same content of Na<sub>2</sub>O, e.g. N9.3S9T40 in comparison with N9.3S0T40, and N9.3S9T60 in comparison with N9.3S0T60. The significantly increased [Al] in the sodium silicate activated fly ash paste was due to the enhanced dissolution of fly ash by the soluble silicate.

516

In the sodium hydroxide activated fly ash paste, the initial preferential dissolution of Al from the 517 fly ash created a siliceous layer on the fly ash grains [57-59]. The initially dissolved aqueous Al 518 species then might absorb to the surface sites on the siliceous layer, passivating the surface by 519 520 preventing the approach of hydroxide ions [60]. Thus, the dissolution of fly ash could be slowed down. In the sodium silicate activated fly ash paste, the soluble silicate could quickly complex 521 the initial preferentially released Al. This prevented Al from absorbing to the surface sites and 522 thus reduced the surface passivation and allowed more rapid dissolution of fly ash [61]. It was 523 reported that the soluble silicate enhanced greater structural breakdown of the frameworks in fly 524 ash if the concentration of Si in the alkaline activator was larger than 213.6 mmol/L [61]. In this 525 study, the concentration of Si in the alkaline activator for the sodium silicate activated fly ash 526 pastes (Table 3) was 4170 mmol/L, twenty times larger than 213.6 mmol/L. As a result, the 527 528 dissolution of Si, Al, Ca, K, Fe and Mg from fly ash was enhanced.

529

The enhanced dissolution of fly ash by soluble silicate can be also supported by the calorimetry 530 data. Figure 10 presents the heat evolution rates of sodium hydroxide activated fly ash and 531 sodium silicate activated fly ash cured at 40 °C and 60 °C. Two calorimetric peaks could be 532 533 identified on the heat evolution rate curves of sodium hydroxide activated fly ash samples cured at 40 °C and 60 °C. The first calorimetric peak (P<sub>1</sub>) corresponds to the dissolution/wetting of fly 534 ash, and the second calorimetric peak (P<sub>2</sub>) corresponds to the acceleration of polymerization 535 536 process of alkali-aluminosilicate type gel [62]. Compared to sodium hydroxide activated fly ash samples, only one peak was identified on the heat evolution rate curves of sodium silicate 537 activated fly ash samples cured at 40 °C and 60 °C. The calorimetry data clearly illustrate the 538 effect of soluble silicate on the dissolution of fly ash. In the sodium hydroxide activated fly ash, 539 the absorbed Al passivated the surface of fly ash and slowed down the dissolution of fly ash. As a 540 result, the acceleration of polymerization of alkali-aluminosilicate type gel was delayed and thus 541 led to the occurrence of the second calorimetric peak. In the sodium silicate activated fly ash, the 542 soluble silicate reduced the surface passivation and allowed rapid dissolution of fly ash. 543 Consequently, the polymerization of alkali-aluminosilicate type gel was accelerated. As such, the 544 calorimetric peak that corresponded to the polymerization was advanced and thus merged into the 545 calorimetric peak that corresponded to the dissolution/wetting of fly ash. For this reason, the 546 second calorimetric peak was not readily identified. 547

548

**Figure 10** Heat evolution rates of sodium hydroxide activated fly ash and sodium silicate activated fly ash cured at 40 °C and 60 °C: (A) N9.3S0T40 in comparison with N9.3S9T40; and (B) N9.3S0T60 in comparison with N9.3S9T60. In the graphs,  $P_1$  and  $P_2$  refer to the first and second calorimetric peak respectively, N and S indicate the

weight percentage of  $Na_2O$  and  $SiO_2$  with respect to fly ash, and T indicates the curing temperature in Celsius. The water to fly ash ratio was 0.35.

554

From Figure 9(C), it is clear that the [OH<sup>-</sup>] decreased with time, and increased significantly with 555 increase of Na<sub>2</sub>O content for both curing temperatures. When sodium silicate was used as the 556 557 alkaline activator, the [OH<sup>-</sup>] decreased significantly in comparison with the sodium hydroxide activated fly ash with the same content of Na<sub>2</sub>O. The decreased alkalinity caused by the increased 558 modulus of alkaline activator  $(SiO_2/Na_2O)$  is in agreement with [63]. The [OH] indicates a pH 559 range of 13.08~14.07 up to 28 days of curing. This pH range shows a higher alkalinity in the pore 560 solution of alkali-activated fly ash than that in the pore solution (pH>12) buffered by Ca(OH)<sub>2</sub> in 561 OPC based materials, and is much higher than the pH of 9.5 below which steel reinforcement 562 corrosion commences [39]. 563

564

The concentration of Ca is plotted in Figure 9(D). The pore solution of sodium silicate activated 565 fly ash paste had higher [Ca] than that of sodium hydroxide activated fly ash paste with the same 566 content of Na<sub>2</sub>O. This can be attributed to the lower alkalinity of pore solution and the enhanced 567 568 dissolution of fly ash by the soluble silicate. The [Ca] decreased with time, which is different from the [Ca] in the pore solutions of alkali-activated slag pastes where no substantial change of 569 the [Ca] was observed. The decrease of [Ca] was due to the precipitation of Ca into solid reaction 570 571 products, which was also observed in [61]. The precipitation of Ca into solid reaction products 572 was further evidenced by the solid phase analysis through XRD. Figure 11 presents the XRD patterns of alkali-activated fly ash samples at 28 days. It can be seen that new crystalline phases 573 574 were formed in addition to quartz (Q), mullite (M) and hematite (H). Chabazite (C) was formed 575 in sodium hydroxide activated fly ash samples, while faujasite (F) was formed in sodium silicate

activated fly ash samples. Both of chabazite and faujasite contain Ca. In addition to the Ca
containing crystalline phases, Ca might be also incorporated into N-A-S-H gels [64]. It has been
reported that Na may be displaced by Ca through ion exchange on the surface of N-A-S-H,
resulting in N-(C-)A-S-H gels [65].

580

Figure 11 X-ray diffraction patterns of alkali-activated fly ash samples cured at 40 °C and 60 °C: (A) N9.3S0T40 and N9.3S0T60; and (B) N9.3S9T40 and N9.3S9T60. In the graphs, Q, M, H, C, F and A refer to quartz, mullite, hematite, chabazite, faujasite and analcime respectively, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to fly ash, and T indicates the curing temperature in Celsius. For all samples, the water to fly ash ratio was 0.35.

586

Figure 9(E) showed an apparent decrease of [Na] with time. When the Na<sub>2</sub>O content increased 587 from 6.2% to 9.3%, the [Na] increased in the sodium hydroxide activated fly ash system. From 588 Figure 9(F), it is clear that the [K] decreased as a function of time for all samples, which is on the 589 contrary to the [K] in the pore solutions of alkali-activated slag pastes (Figure 3(F)). In the 590 sodium hydroxide/silicate activated fly ash based materials, sodium based crystalline phases are 591 normally identified, such as Na-chabazite and hydroxysodalite [66, 67]. On the contrary, 592 potassium based crystalline phases are rarely reported. This is due to the better zeolitization 593 594 capabilities of sodium cations than those of potassium cations since the sodium cations are smaller than the potassium cations [68]. So, the decrease of [K] was most likely to be attributed 595 to the formation of potassium-aluminosilicate (K-A-S-H) type gels. The formation of K-A-S-H in 596 alkali-activated fly ash has been reported in the literature [69, 70]. The increase of Na<sub>2</sub>O content 597 from 6.2% to 9.3% did not show much influence on [K]. Comparing the sodium silicate activated 598 fly ash paste to the sodium hydroxide activated fly ash paste with the same content of Na<sub>2</sub>O, the 599

600 [K] increased about two times. This could be due to the enhanced dissolution of fly ash by the 601 soluble silicate.

602

The concentrations of Fe and Mg in the pore solutions of alkali-activated fly ash pastes are plotted as a function of time in Figure 9(G) and Figure 9(H) respectively (hereafter denoted as [Fe] and [Mg] respectively). It is clear that the concentrations of Fe and Mg decreased with time. Since the dissolution of fly ash is enhanced in the sodium silicate activated fly ash paste, the concentrations of Fe and Mg were higher when compared to those in the sodium hydroxide activated fly ash paste with the same content of Na<sub>2</sub>O.

609

Figure 9(I) plots the concentration of S as a function of time (hereafter denoted as [S]). The [S] 610 increased with time for all alkali-activated fly ash samples. For sodium hydroxide activated fly 611 ash, the [S] increased with increase of Na<sub>2</sub>O content. Fly ash contains 0.73% of SO<sub>3</sub> (see Table 612 1). If the S in the fly ash is completely dissolved in the alkaline activator, the concentration of S 613 is around 250 mmol/L. However, the [S] measured in the pore solution was around 350 mmol/L, 614 615 much larger than 250 mmol/L. This was because of the decrease of the solution volume resulted from the dissolution/wetting of fly ash and the polymerization process of alkali-aluminosilicate 616 type gel. It is known that the porosity of alkali-activated fly ash decreases with time [28]. This 617 indicates the decrease of the volume of solution in the alkali-activated fly ash. Therefore, using 618 the volume of alkaline activator to calculate the concentration of S would result in 619 underestimation. 620

621

622 3.4 Pore solution composition of alkali-activated slag blended with fly ash paste

To study the influence of the replacement of slag by fly ash on the pore solution composition, one 624 625 mixture of alkali-activated slag blended with fly ash was used in this study. Figure 12(A) presents the elemental concentrations in the pore solution of alkali-activated slag blended with fly ash 626 (N6S5.4B). The pore solution data are listed in the Appendix. It can be seen that [Si], [Na] and 627 628 [OH<sup>-</sup>] decreased as a function of time, while [Al] and [Ca] did not show much change with time. The [OH] indicates a pH range of 14.05~14.37 up to 28 days. Compared to the alkali-activated 629 slag (N6S5.4) with the same contents of Na<sub>2</sub>O and SiO<sub>2</sub> as shown in Figure 12(B), 50 wt.% 630 replacement of slag by fly ash did not result in a substantial change on the pore solution 631 composition of alkali-activated slag blended with fly ash paste. 632

633

Figure 12 Elemental concentrations in the pore solutions of alkali-activated slag blended with fly ash paste (A), and
alkali-activated slag paste (B). In the graphs, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect
to the precursor (slag, or blend of slag and fly ash), and B indicates blend of slag and fly ash. The curing temperature
was 20 °C. The water to precursor ratio was 0.4.

638

But at a later curing age, the pore solution of alkali-activated slag blended with fly ash paste had 639 640 higher [Si] and lower [Na] than that of alkali-activated slag paste with the same alkaline activator. At 28 days (672 hours), the concentrations of Si and Na were 37.5 mmol/L and 1670.4 641 mmol/L respectively in the pore solution of alkali-activated slag blended with fly ash paste, while 642 they were 11.9 mmol/L and 2517.7 mmol/L respectively in the pore solution of alkali-activated 643 644 slag paste. The higher [Si] in the alkali-activated slag blended with fly ash paste was because fly ash contains more reactive SiO<sub>2</sub> (37%) than slag (32.9%). The lower [Na] in the alkali-activated 645 slag blended with fly ash paste was because the replacement of slag by fly ash led to the 646 formation of alkali-aluminosilicate type gel. The alkali-aluminosilicate type gel was reported to 647

be formed together with calcium aluminosilicate type gel in alkali-activated blend of slag and fly
ash in the literature [64, 71, 72]. The co-existence of calcium-aluminosilicate and alkalialuminosilicate type gels was also confirmed by the solid phase analysis through XRD and FTIR.

652 Figure 13(A) presents the XRD patterns of alkali-activated slag blended with fly ash and alkaliactivated slag with the same alkaline activator, e.g. N6S5.4B in comparison with N6S5.4 at 28 653 days. Like in alkali-activated slag paste (N6S5.4), calcium aluminosilicate hydrate was also 654 formed in alkali-activated slag blended with fly ash paste. Since alkali-aluminosilicate type gel is 655 amorphous to semi-crystalline [73], it was not detected by XRD. As such, FTIR technique was 656 used to investigate the bonding environments in non-crystalline phases. Figure 13(B) plots the 657 FTIR spectra of N6S5.4B and N6S5.4 at 28 days. The overall broad band of interest was located 658 between 800 and 1200 cm<sup>-1</sup>. This band is associated with the asymmetric stretching mode of Si-659 660 O-T bonds (T: tetrahedral Si or Al), and is usually used to study the changes of the amorphous gel structure [74, 75]. The Si-O-T band was centered at 948 cm<sup>-1</sup> in alkali-activated slag paste, 661 while it was centered at 952 cm<sup>-1</sup> in alkali-activated slag blended with fly ash paste. The shift of 662 Si-O-T band towards to a higher wavenumber was attributed to the incorporation of fly ash that 663 resulted in the formation of more crosslinked alkali-aluminosilicate type gel [64]. As such, it can 664 be concluded that calcium aluminosilicate and alkali-aluminosilicate type gels were both formed 665 in alkali-activated slag blended with fly ash paste (N6S5.4B). 666

667

Figure 13 Solid phase analysis of alkali-activated slag blended with fly ash and alkali-activated slag with the same
alkaline activator: (A) X-ray patterns, and (B) FTIR spectra. In the graphs, Q, M, H, Ht, K and C-(N-)A-S-H refer to
quartz, mullite, hematite, hydrotalcite, katoite and calcium aluminosilicate hydrate respectively, N and S indicate the

weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to the precursor (slag, or blend of slag and fly ash), and B indicates
blend of slag and fly ash. The curing temperature was 20 °C. The water to precursor ratio was 0.4.

673

Compared to the calcium aluminosilicate hydrate type gel, the alkali-aluminosilicate type gel 674 takes up a larger amount of Na to balance the charge of Al in the bonding network [76]. As such, 675 676 the pore solution of alkali-activated slag blended with fly ash paste had a relatively lower concentration of Na than that of alkali-activated slag paste with the same alkaline activator. The 677 leaching of Na from AAMs is driven by the concentration gradients between the pore solution 678 and the water that surrounds the materials [26]. From this point of view, the decreased 679 concentration of Na in the pore solution due to the replacement of slag by fly ash can lead to 680 decreases of the concentration gradients and thus reduce the leaching of Na in the alkali-activated 681 slag blended with fly ash paste. 682

683

684 3.5 Conceptual models to describe the elemental concentrations in the pore solutions of alkali-685 activated slag/fly ash pastes

686

687 Figure 14(A) presents the conceptual model to describe the concentrations of Al, Ca, K, Fe, and Mg. Since Al, Ca, K, Fe and Mg are sourced from the precursor, the concentrations of these 688 elements in the pore solution initially start from zero and increase with the progressive 689 690 dissolution of the precursor. When the concentrations of these elements increase to be 691 oversaturated with respect to solids, these elements are thermodynamically favored to precipitate and form the solids. If there are nuclei of the solids formed and the nuclei reach a critical size, the 692 693 solids start to grow. The formation of nuclei and the growth of the solids consume the elements 694 that constitute the solids. In the meantime, the dissolution of the precursor continues, releasing

these elements into the pore solution. When the consumption rates of these elements exceed the release rates of these elements, the concentrations of these elements decrease with time. Since the continuous release of these elements can partially compensate the consumption of these elements, the concentrations of these elements keep oversaturated with respect to the solids with time. But the degrees of oversaturation with respect to the solids decrease because the consumption of these elements cannot be completely replenished by the dissolution of the precursor.

701

702 Since Si can be sourced from both the alkaline activator and the dissolution of the precursor, the conceptual model to describe the concentration of Si in the pore solution depends on the 703 concentration of Si in the alkaline activator ([Si]<sub>0</sub>). If [Si]<sub>0</sub>=0 (Figure 14(B)), Si can only be 704 sourced from the dissolution of the precursor and its concentration in the pore solution initially 705 starts from zero and increase with the progressive dissolution of the precursor. When the release 706 rate of Si is equal to the consumption rate of Si, the concentration of Si reaches the maximum 707 708 concentration ( $[Si]_{max}$ ), after which it decreases with time. If  $0 < [Si]_0 < [Si]_{max}$  (Figure 14(C)), the concentration of Si in the pore solution initially starts from [Si]<sub>0</sub>. Since [Si]<sub>0</sub><[Si]<sub>max</sub>, the release 709 710 rate of Si from the precursor is larger than the consumption rate of Si. So, the concentration of Si increases with the progressive dissolution of the precursor until it reaches [Si]<sub>max</sub>, at this point the 711 release rate of Si from fly ash is equal to the consumption rate of Si, after which it decreases with 712 time. If  $[Si]_0 \ge [Si]_{max}$  (Figure 14(D)), the concentration of Si in the pore solution initially starts 713 from  $[Si]_0$  and decreases with time because the  $[Si]_0$  enables a consumption rate that is not 714 715 smaller than the release rate of Si.

Figure 14 Conceptual models to describe the concentrations of Al, Ca, K, Fe, Mg, Si, Na, OH<sup>-</sup> and S in the pore
solution of alkali-activated slag/fly ash paste. In the graphs, [Si]<sub>0</sub> is the concentration of Si in the alkaline activator.
[Si]<sub>max</sub> is the maximum concentration of Si.

720

Figure 14(E) shows the conceptual model for the description of the concentrations of Na and OH<sup>-</sup> 721 . Besides trace amount of Na<sub>2</sub>O in the precursors, Na in the pore solution is sourced from the 722 alkaline activator. With the continuous formation of solids, for example the alkali-aluminosilicate 723 type gel, the Na is consumed and thus the concentration of Na decreases with time. Although the 724 dissolution of alkali-earth metals (Ca, Mg etc.) from the precursor may result in formation of 725 hydroxyl ion in the pore solution, this amount of hydroxyl ions are much smaller than the amount 726 consumed by the dissolution of Si and Al. So, the concentration of OH decreases with time. S is 727 728 sourced from slag/fly ash. Since no S related solids precipitate (as no S containing phases 729 identified, see Figures 8, 11 and 13(A)), the concentration of S in the pore solution increases with 730 time and reaches a maximum when S is completely dissolved from the precursor (Figure 14(F)). 731

3.6 The influence of temperature on the pore solution composition of alkali-activated fly ashpaste

734

The increase of temperature led to decreases of the elemental concentrations in the pore solutions of alkali-activated fly ash pastes (see Figure 9). The influence of temperature on the pore solution composition of alkali-activated fly ash pastes can be discussed from the aspects of kinetics and thermodynamics.

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740 3.6.1 Kinetics

741

The dissolution of fly ash releases the elements into the pore solution, while the formation of solid reaction products consumes the elements that constitute the solid reaction products. So, the elemental concentrations in the pore solution depend on the competition between the dissolution of fly ash and the formation of solid reaction products. The increase of temperature leads to accelerations of both the dissolution of fly ash and the formation of solid reaction products [77, 78]. It means the increase of temperature accelerates both the release of elements from fly ash and the consumption of these elements by the formation of solid reaction products.

749

Figure 15 plots the heat evolution rates of sodium hydroxide activated fly ash pastes at 40 °C and 750 60 °C. The calorimetric responses of sodium silicate activated fly ash pastes show only one 751 calorimetric peak (see Figure 10). This makes it impossible to separate the calorimetric response 752 that corresponds to the dissolution of fly ash, and the calorimetric response that corresponds to 753 the formation of alkali-aluminosilicate type gel. Thus, these results are not included in Figure 15. 754 Comparing the calorimetric responses at 60 °C to the calorimetric responses at 40 °C, it can be 755 756 seen that the increase of temperature led to a larger increase of the intensity of the second calorimetric peak. This indicates the increase of temperature had a larger acceleration of the 757 formation of solid reaction products than that of the dissolution of fly ash. 758

759

Figure 15 Heat evolution rates of alkali-activated fly ash pastes at 40 °C (A) and 60 °C (B). In the graphs, P<sub>1</sub> and P<sub>2</sub>
refer to the first and second calorimetric peak respectively, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub>
with respect to fly ash, and T indicates the curing temperature in Celsius. The water to fly ash ratio was 0.35.

Since the increase of temperature led to a larger acceleration of the consumption of elements than 764 765 that of the release of elements, the consumption rates of elements exceed the release rates of 766 elements at lower elemental concentrations. So, maximum elemental concentrations after the increase of temperature were smaller than the maximum concentrations before the increase of 767 768 temperature. To show the influence of temperature on the pore solution composition of alkaliactivated fly ash paste. Figure 16 plots the concentrations of Al. Ca. K. Fe and Mg before and 769 after the increase of temperature. In the experiments, the alkali-activated fly ash pastes were in 770 771 fluid state at the early curing time, which made it infeasible to collect the pore solution using the steel-die method. For this reason, the maximum elemental concentrations were not obtained. As 772 shown in Figure 16, the experimental measurements started after the maximum elemental 773 774 concentrations.

775

Figure 16 The influence of an increase of temperature on the concentrations of Al, Ca, K, Fe and Mg in the pore
solution of alkali-activated fly ash paste. T<sub>1</sub> and T<sub>2</sub> are temperatures.

778

#### 779 3.6.2 Thermodynamics

780

Thermodynamics assumes a solubility equilibrium between the solid and the aqueous species in the liquid that react to form the solid [17]. From this point of view, the solubility of the reaction products also influence the concentrations of the elements that react to form the solids. The alkali-aluminosilicate type gel, as the main solid reaction products in alkali-activated fly ash paste [79], is amorphous to semi-crystalline [73] and considered as the precursors to zeolites in the literature [75, 80]. It is reported that an increase of temperature leads to increase of longrange ordering and higher degree of crosslinking in alkali-aluminosilicate type gel [78, 81, 82].

This point can be further confirmed by the FTIR analysis. Figure 17 presents the infrared spectra 788 of alkali-activated fly ash samples at 28 days. It can be observed that an increase of temperature 789 790 (from 40 °C to 60 °C) led to the shift of the Si-O-T band to higher wavenumbers, from 971 to 980 cm<sup>-1</sup> for sodium hydroxide activated fly ash and from 979 to 983 cm<sup>-1</sup> for sodium silicate 791 792 activated fly ash. The shift of Si-O-T band from a lower wavenumber to a higher wavenumber indicates a higher degree of crosslinking of alkali-aluminosilicate type gel [64] and that the 793 structure of alkali-aluminosilicate type gel becomes more ordered and stable [75]. As such, an 794 increase of temperature leads to a higher degree of crosslinking of alkali-aluminosilicate type gel 795 and a more stable structure. 796

797

Figure 17 FTIR spectra of alkali-activated fly ash samples at 28 days. In the graph, N and S indicate the weight
percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to fly ash, and T indicates the curing temperature in Celsius. The water to
fly ash ratio was 0.35.

801

The higher degree of crosslinking and more stable structure of alkali-aluminosilicate type gel could lead to a decrease of the solubility of alkali-aluminosilicate type gel. The decreased solubility of alkali-aluminosilicate type gel results in lower concentrations of the elements that react to form the alkali-aluminosilicate type gel. So, the increase of temperature from 40 °C to 60 °C led to decreases of the concentrations of Si, Al, Na and K that reacted to form alkali aluminosilicate type gel.

808

```
809 4 Conclusions
```

This study investigated the pore solution composition of alkali-activated slag/fly ash pastes by means of ICP-OES analysis technique. The following conclusions can be drawn based on this study:

814

- (1) In preparation of the samples for the ICP-OES analysis, alkaline dilution should be used
  for determination of the concentration of S. The acid dilution results in evaporation of
  H<sub>2</sub>S and white precipitation of S. In the pore solutions of alkali-activated slag/fly ash
  pastes, the elemental concentrations depend on the alkaline activator and curing
  temperature.
- 820

(2) The pore solutions of alkali-activated slag pastes are oversaturated with respect to solids.
In the initial period, the effective saturation indexes with respect to calcium
aluminosilicate hydrates are relatively high and decrease with time. On the contrary, the
effective saturation indexes with respect to the crystalline phases increase rapidly. After
the initial period, the effective saturation indexes show no substantial change with time.

- 826
- (3) In the pore solutions of alkali-activated fly ash pastes, an increase of temperature leads to
  decreases of the concentrations of Si, Al, Ca, K, Fe and Mg, while the soluble silicate in
  the alkaline activator results in increases of these elemental concentrations.
- 830
- (4) Compared to the alkali-activated slag paste with the same alkaline activator, 50 wt.%
  replacement of slag by fly ash does not result in a substantial influence on the pore
  solution composition of alkali-activated slag blended with fly ash paste.
- 834
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(5) According to the experimental results, conceptual models are proposed to describe the
elemental concentrations in the pore solutions. Based on the conceptual models, the
influence of temperature on the pore solution composition of alkali-activated fly ash paste
can be well interpreted. In addition, these conceptual models will help to develop kinetic
models to describe quantitatively the developments of the elemental concentrations with
time.

841

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843

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848

### 849 Appendix. Pore solution composition data of alkali-activated slag/fly ash pastes

850

The pore solution data, i.e. elemental concentrations of Si, Al, OH<sup>-</sup>, Ca, Na, K, Fe, Mg and S, are
listed in Tables A1-A9.

853

**Table A1** Concentration of Si (mmol/L)

855

**Table A2** Concentration of A1 (mmol/L)

857

858 **Table A3** Concentration of OH<sup>-</sup> (mmol/L)

859	
860	Table A4 Concentration of Ca (mmol/L)
861	
862	Table A5 Concentration of Na (mmol/L)
863	
864	Table A6 Concentration of K (mmol/L)
865	
866	Table A7 Concentration of Fe (mmol/L)
867	
868	Table A8 Concentration of Mg (mmol/L)
869	
870	Table A9 Concentration of S (mmol/L)
871	
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# 1134 Tables

1135

**Table 1** Chemical compositions of blast furnace slag and fly ash (by weight, %)

	Oxide	SiO <sub>2</sub>	CaO	$Al_2O_3$	MgO	Fe <sub>2</sub> O <sub>3</sub>	SO <sub>3</sub>	K <sub>2</sub> O	TiO <sub>2</sub>	L.I.*
	Slag	32.91	40.96	11.85	9.23	0.46	1.61	0.33	1.00	1.15
	Fly ash	52.90	4.36	26.96	1.50	6.60	0.73	1.31	1.14	3.37
1137	*L.I. refe	rs to loss o	n ignition.							
1138										
1139										
1140										
1141	Table 2	Mixture co	mpositions a	and curing to	emperature					

Sample <sup>a</sup>	Precursor	$Na_2O^b$	SiO <sub>2</sub> <sup>b</sup>	Curing temperature
N4S0	slag	4	0	20 °C
N6S0	slag	6	0	20 °C
N8S0	slag	8	0	20 °C
N4S2.7	slag	4	2.7	20 °C
N4S5.4	slag	4	5.4	20 °C
N6S5.4	slag	6	5.4	20 °C
N8S5.4	slag	8	5.4	20 °C
N6.2S0T40	fly ash	6.2	0	40 °C
N9.3S0T40	fly ash	9.3	0	40 °C
N9.3S9T40	fly ash	9.3	9	40 °C
N6.2S0T60	fly ash	6.2	0	60 °C
N9.3S0T60	fly ash	9.3	0	60 °C
N9.3S9T60	fly ash	9.3	9	60 °C
N6S5.4B	blend <sup>c</sup>	6	5.4	20 °C

- <sup>a</sup>N and S indicates weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to precursor, T indicates temperature in Celsius
- and B indicates blend of slag and fly ash.
- <sup>b</sup>weight percentage with respect to precursor content (wt. %).
- 1145  $^{c}$  blend = 50 wt. % slag + 50 wt. % fly ash.
- 1146
- 1147
- 1148
- 1149
- 1150
- 1151

Sample <sup>a</sup>	pН	[Na] (mmol/L)	[Si] (mmol/L)	[OH <sup>-</sup> ] (mmol/L) <sup>b</sup>
N4S0	14.31	3239	0	3239(1957)
N6S0	14.50	4857	0	4857(2517)
N8S0	14.64	6466	0	6466(2953)
N4S2.7	14.14	3204	1117	1757(831)
N4S5.4	13.87	3193	2227	821(261)
N6S5.4 and N6S5.4B	14.30	4773	2219	2218(578)
N8S5.4	14.52	6339	2211	3636(817)
N6.2S0T40 and N6.2S0T60	14.58	5731	0	5731(2766)
N9.3S0T40 and N9.3S0T60	14.79	8564	0	8564(3408)
N9.3S9T40 and N9.3S9T60	14.60	8340	4170	3908(257)

1152 Table 3 The pH and concentrations of Na, Si, and OH<sup>-</sup> in the alkaline activators

<sup>a</sup>N and S indicates weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to precursor, T indicates temperature in Celsius

<sup>b</sup>The numbers outside the brackets were obtained by thermodynamic calculations including the hydroxide from

- 1156 NaOH(aq), while those inside the brackets were obtained by excluding the hydroxide from NaOH(aq).
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- 1158

1159 Table 4 Chemical reactions and equilibrium solubility products at 25°C and 1 bar for eight end-members and

1160 crystalline reaction products in alkali-activated slag

End-member	Chemical reactions	Log K <sub>s0</sub>
C- $(N$ - $)A$ - $S$ - $Hg$	el ideal solid solution eight end-members, 'CNASH_ss' model [36]	
5CA	$(CaO)_{1.25} \cdot (Al_2O_3)_{0.125} \cdot (SiO_2) \cdot (H_2O)_{1.625}$	-10.75
	$\Leftrightarrow 1.25 \text{Ca}^{2+} + \text{SiO}_3^{2-} + 0.25 \text{AlO}_2^{-} + 0.25 \text{OH}^{-} + 1.5 \text{H}_2 \text{O}$	
INFCA	$(CaO) \cdot (Al_2O_3)_{0.15625} \cdot (SiO_2)_{1.1875} \cdot (H_2O)_{1.65625} + 0.68750 H^-$	-8.90
	$\Leftrightarrow \text{ Ca}^{2+} + 1.1875\text{SiO}_3^{2-} + 0.3125\text{AlO}_2^{-} + 2\text{H}_2\text{O}$	
5CNA	$(CaO)_{1.25} \cdot (Na_2O)_{0.25} \cdot (Al_2O_3)_{0.125} \cdot (SiO_2) \cdot (H_2O)_{1.25}$	-10.40
	$\Leftrightarrow 1.25Ca^{2+} + SiO_3^{2-} + 0.25AlO_2^{-} + 0.5Na^{+} + 0.75OH^{-} + H_2O$	
INFCNA	$(CaO) \cdot (Na_2O)_{0.34375} \cdot (Al_2O_3)_{0.15625} \cdot (SiO_2)_{1.1875} \cdot (H_2O)_{1.3}$	-10.00
	$\Leftrightarrow \text{ Ca}^{2+} + 1.1875\text{Si}0_3^{2-} + 0.3125\text{Al}0_2^{-} + 0.6875\text{Na}^{+} + 1.3125\text{H}_2\text{O}$	
INFCN	$(CaO) \cdot (Na_2O)_{0.3125} \cdot (SiO_2)_{1.5} \cdot (H_2O)_{1.1875} + 0.3750H^-$	-10.70
	$\Leftrightarrow \text{ Ca}^{2+} + 1.5\text{Si}0_3^{2-} + 0.625\text{Na}^+ + 1.375\text{H}_2\text{O}$	
T2C*	$(CaO)_{1.5} \cdot (SiO_2) \cdot (H_2O)_{2.5} \iff 1.5Ca^{2+} + SiO_3^{2-} + OH^- + 2H_2O$	-11.60
T5C*	$(CaO)_{1.25} \cdot (SiO_2)_{1.25} \cdot (H_2O)_{2.} \Leftrightarrow 1.25Ca^{2+} + 1.25SiO_3^{2-} + 2.5H_2O$	-10.50
TobH*	$(\text{CaO}) \cdot (\text{SiO}_2)_{1.5} \cdot (\text{H}_2\text{O})_{2.5} + 0\text{H}^- \Leftrightarrow \text{Ca}^{2+} + 1.5\text{SiO}_3^{2-} + 3\text{H}_2\text{O}$	-7.90
Crystalline rea	action products in alkali-activated slag [17, 33]	
C <sub>2</sub> ASH <sub>8</sub>	$(CaO)_2 \cdot (Al_2O_3) \cdot (SiO_2) \cdot (H_2O)_8 \iff 2Ca^{2+} + 2AlO_2^- + SiO_3^{2-} + 8H_2O$	-19.10
C <sub>3</sub> AH <sub>6</sub>	$(CaO)_3 \cdot (Al_2O_3) \cdot (H_2O)_6 \Leftrightarrow 3Ca^{2+} + 2AlO_2^- + 4OH^- + 4H_2O$	-20.85
Ca(OH) <sub>2</sub>	$Ca(OH)_2 \Leftrightarrow Ca^{2+} + 2OH^-$	-5.20
$C_4AH_{13}$	$(CaO)_4 \cdot (Al_2O_3) \cdot (H_2O)_{13} \Leftrightarrow 4Ca^{2+} + 2AlO_2^- + 6OH^- + 10H_2O$	-25.41

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and B indicates blend of slag and fly ash.

Comula	Time (hours)									
Sample	3	7	24	72	168	336	672			
N4S0	24.9	17.9	9.6	5.9	4.6	4.5	3.7			
N6S0	39.7	33.7	22.3	14.3	11.5	9.1	7.7			
N8S0	44.6	45.7	42.2	27.4	21.4	18.4	16.3			
N4S2.7	<sup>a</sup>		13.6	7.6	6.1	5.5	5.1			
N4S5.4	469.6	312.9	22.4	12.1	9.7	7.5	6.4			
N6S5.4	172.6	40.9	29.0	18.4	15.6	13.6	11.9			
N8S5.4	78.9	67.1	50.9	38.2	30.9	28.6	25.7			
N6.2S0T40				990.9	922.5		219.1			
N9.3S0T40				2123.4	1048.0		150.2			
N9.3S9T40				1965.9	2089.6		1381.3			
N6.2S0T60			1107.3	374.3	113.6		12.8			
N9.3S0T60			1504.5	552.1	221.6		74			
N9.3S9T60			2421.6	2043.6	1787.7		655.6			
N6S5.4B		227.9	63.8	56.5	37.5		37.5			

### 1164 Table A1 Concentration of Si (mmol/L)

<sup>a</sup>-- means not measured.

#### 1166

## **Table A2** Concentration of Al (mmol/L)

	Time (hours)								
Sample	3	7	24	72	168	336	672		
N4S0	10.3	10.2	9.7	8.8	8.4	7.9	7.5		
N6S0	15.4	15.4	15.1	14.6	14.2	12.4	12.1		
N8S0	24.4	23.3	22.9	21.6	20.4	19.5	19.1		
N4S2.7	<sup>a</sup>		15.7	15.4	14.4	14.4	13.3		
N4S5.4	11.8	10.8	9.7	12.2	13.2	12.4	11.1		
N6S5.4	17.3	23.3	22.3	22.1	21.7	20.6	18.9		
N8S5.4	32.7	32.3	32.9	32.3	30.7	31.3	29.3		
N6.2S0T40				41.7	22.7		2.0		
N9.3S0T40				177.8	32.8		1.1		
N9.3S9T40				207.9	144.0		12.8		
N6.2S0T60			26.9	3.1	0.3				
N9.3S0T60		-	35.8	4.4	1.0		0.3		
N9.3S9T60			128.0	36.9	15.9		2.3		
N6S5.4B		21.4	30.9	29.7	30.1		25.4		

 $( \mathbf{N} )$ 

<sup>a</sup>-- means not measured.

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Commlo	Time (hours)								
Sample	3	7	24	72	168	336	672		
N4S0	2098	2153	1890	1667	1581	1410	1289		
N6S0	2907	3145	2864	2640	2537	2468	2213		
N8S0	4149	4076	3989	3910	3883	3572	3621		
N4S2.7	<sup>a</sup>		1372	1070	1100	1009	969		
N4S5.4			833	783	803	702	589		
N6S5.4	2251	2074	1882	1747	1550	1426	1270		
N8S5.4	2926	2941	2695	2609	2478	2406	2163		
N6.2S0T40				2525	1015		333		
N9.3S0T40					2327		1177		
N9.3S9T40					1292		424		
N6.2S0T60			1223	432	218		119		
N9.3S0T60			2295	1643	1290		628		
N9.3S9T60			1486	610	250		202		
N6S5.4B		2370					1128		

### **1176** Table A3 Concentration of OH<sup>-</sup> (mmol/L)

 $\overline{a}$ -- means not measured.

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## **Table A4** Concentration of Ca (mmol/L)

Sampla	Time (hours)									
Sample	3	7	24	72	168	336	672			
N4S0	0.64	0.58	0.63	0.66	0.67	0.81	0.76			
N6S0	0.53	0.47	0.36	0.53	0.58	0.52	0.57			
N8S0	0.59	0.56	0.51	0.44	0.59	0.56	0.54			
N4S2.7	<sup>a</sup>		0.32	0.32	0.34	0.35	0.39			
N4S5.4	2.34		1.49	1.40	1.24	1.28	1.56			
N6S5.4	0.58	0.46	0.51	0.49	0.60	0.65	0.75			
N8S5.4	0.50	0.40	0.35	0.35	0.36	0.26	0.23			
N6.2S0T40				4.05	2.83		1.52			
N9.3S0T40				12.23	3.80		1.10			
N9.3S9T40				17.23	16.1		1.67			
N6.2S0T60			2.33	1.10	0.69		0.99			
N9.3S0T60			2.70	0.98	0.61		0.71			
N9.3S9T60			11.18	3.93	1.78		0.98			
N6S5.4B		0.71	1.16	0.69	0.56		1.02			

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<sup>a</sup>-- means not measured.

Sample	Time (hours)						
	3	7	24	72	168	336	672
N4S0	2793	2729	2459	2184	2141	1948	1747
N6S0	4064	4231	3956	3606	3518	3467	3077
N8S0	6291	5542	5948	5722	5162	5374	5207
N4S2.7	<sup>a</sup>		1994	1879	1862	1802	1735
N4S5.4	2436	2254	1426	1386	1508	1387	1343
N6S5.4	3252	3136	3007	2750	2664	2600	2518
N8S5.4	4370	4322	4160	4104	4149	4122	3774
N6.2S0T40				3990	2268		1323
N9.3S0T40				5287	3191		1680
N9.3S9T40				4450	3802		2155
N6.2S0T60			2489	1335	1027		792
N9.3S0T60			3117	2071	1651		1443
N9.3S9T60			3669	2612	2395		1631
N6S5.4B		3921	3189	2804	2515		1670

#### 1188 Table A5 Concentration of Na (mmol/L)

<sup>a</sup>-- means not measured.

#### 1190

## **Table A6** Concentration of K (mmol/L)

Sampla	Time (hours)							
Sample	3	7	24	72	168	336	672	
N4S0	30.7	39.7	52.6	61.7	72.2	77.7	81.2	
N6S0	40.6	53.6	65.4	83.0	98.6	109.8	124.5	
N8S0	57.9	60.6	72.4	87.4	105.4	120.9	139.9	
N4S2.7	<sup>a</sup>		33.0	44.3	48.4	54.6	56.9	
N4S5.4	7.6	8.5	15.7	26.0	33.6	32.8	35.4	
N6S5.4	16.5	41.1	45.9	56.1	62.3	67.0	62.2	
N8S5.4	52.0	56.7	68.1	78.3	88.5	94.9	93.2	
N6.2S0T40				88.4	61.5		26.4	
N9.3S0T40				81.9	49.0		24.9	
N9.3S9T40				126.4	115.4		50.8	
N6.2S0T60			56.6	22.5	14.8		9.8	
N9.3S0T60		-	39.6	23.7	16.1		13.5	
N9.3S9T60			104.0	59.5	49.9		24.4	
N6S5.4B		20.0	37.2	46.4	50.7		44.1	

<sup>a</sup>-- means not measured.

1193

# **Table A7** Concentration of Fe (mmol/L)

Samula		Time	(hours)	
Sample	24	72	168	672
N6.2S0T40	<sup>a</sup>	1.72	1.11	0.12
N9.3S0T40		9.05	1.49	
N9.3S9T40		12.46	8.64	0.32
N6.2S0T60	0.88	0.15		
N9.3S0T60	1.30	0.13		
N9.3S9T60	4.375	1.55	0.52	0.20

1195 <sup>a</sup>-- means not measured.

#### 1197 Table A8 Concentration of Mg (mmol/L)

Samula		Time	e (hours)	
Sample	24	72	168	672
N6.2S0T40	<sup>a</sup>	1.58	1.29	0.19
N9.3S0T40		8.04	1.19	0.09
N9.3S9T40		9.29	7.92	0.39
N6.2S0T60	0.75	0.13		
N9.3S0T60	0.92	0.11		0.06
N9.3S9T60	3.83	1.21	1.42	0.18

<sup>a</sup>-- means not measured.

1199

1200 Table A9 Concentration of S (mmol/L)

	Time (hours)					
Sample	24	72	168	672		
N6.2S0T40	<sup>a</sup>	253	274	356		
N9.3S0T40		345	367	449		
N9.3S9T40		40	47	-63		
N6.2S0T60	238	287	339	317		
N9.3S0T60	315	360	371	391		
N9.3S9T60	20	25	25	40		

1201 <sup>a</sup>-- means not measured.

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# 1216 Figures



Figure 1 X-ray diffraction patterns of slag and fly ash. In the graph, Q, M and H refer to quartz, mullite and hematite respectively.



Figure 2 White precipitation and its element composition in diluted pore solution using nitric acid.



Figure 3 Elemental concentrations in the pore solutions of alkali-activated slag pastes. In the graphs, N and S indicate weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The water to slag ratio was 0.4 and the curing temperature was 20 °C.



**Figure 4** Concentrations of one element as a function of the concentration of another element in pore solution. (A): [Na], [Al] and [Ca] versus [OH<sup>-</sup>]. (B): [Si] versus [OH<sup>-</sup>]. (C) [Si] versus [Ca]. In the graphs, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The curing temperature was 20 °C. The water to slag ratio was 0.4.





Figure 5 The ionic strengths in the pore solutions of alkali-activated slag pastes, calculated by GEM-Selektor. In the graphs, N and S indicate the weight percentage of  $Na_2O$  and  $SiO_2$  with respect to slag. The curing temperature was 20 °C. The water to slag ratio was 0.4.





**Figure 6** Effective saturation indexes with respect to the eight end-members as functions of time. In the graphs, N and S indicate the weight percentage of  $Na_2O$  and  $SiO_2$  with respect to slag. The curing temperature was 20 °C. The water to slag ratio was 0.4. A ESI of 0 indicates equilibrium between solution and solid; a positive ESI indicates oversaturation and a negative ESI indicates undersaturation.





**Figure 7** Effective saturation indexes with respect to  $C_2ASH_8$ ,  $C_3AH_6$ ,  $C_4AH_{13}$  and  $Ca(OH)_2$  as functions of time. In the graphs, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to slag. The curing temperature was 20 °C. The water to slag ratio was 0.4. A ESI of 0 indicates equilibrium between solution and solid; a positive ESI indicates oversaturation and a negative ESI indicates undersaturation.



Figure 8 X-ray diffraction patterns of N4S0, N8S0 and N4S5.4 at 28 days. In the graphs, N and S indicate the weight percentage of  $Na_2O$  and  $SiO_2$  with respect to slag. The curing temperature was 20 °C. The water to slag ratio was 0.4.





Figure 9 Elemental concentrations in the pore solutions of alkali-activated fly ash pastes where N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to fly ash, and T indicates the curing temperature in Celsius. The water to fly ash ratio was 0.35.





**Figure 10** Heat evolution rates of sodium hydroxide activated fly ash and sodium silicate activated fly ash cured at 40 °C and 60 °C: (A) N9.3S0T40 in comparison with N9.3S9T40; and (B) N9.3S0T60 in comparison with N9.3S9T60. In the graphs,  $P_1$  and  $P_2$  refer to the first and second calorimetric peak respectively, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to fly ash, and T indicates the curing temperature in Celsius. The water to fly ash ratio was 0.35.

- 1232
- 1233
- 1234



**Figure 11** X-ray diffraction patterns of alkali-activated fly ash samples cured at 40 °C and 60 °C: (A) N9.3S0T40 and N9.3S0T60; and (B) N9.3S9T40 and N9.3S9T60. In the graphs, Q, M, H, C, F and A refer to quartz, mullite, hematite, chabazite, faujasite and analcime respectively, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to fly ash, and T indicates the curing temperature in Celsius. For all samples, the water to fly ash ratio was 0.35.



**Figure 12** Elemental concentrations in the pore solutions of alkali-activated slag blended with fly ash paste (A), and alkali-activated slag paste (B). In the graphs, N and S indicate the weight percentage of  $Na_2O$  and  $SiO_2$  with respect to the precursor (slag, or blend of slag and fly ash), and B indicates blend of slag and fly ash. The curing temperature was 20 °C. The water to precursor ratio was 0.4.



**Figure 13** Solid phase analysis of alkali-activated slag blended with fly ash and alkali-activated slag with the same alkaline activator: (A) X-ray patterns, and (B) FTIR spectra. In the graphs, Q, M, H, Ht, K and C-(N-)A-S-H refer to quartz, mullite, hematite, hydrotalcite, katoite and calcium aluminosilicate hydrate respectively, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to the precursor (slag, or blend of slag and fly ash), and B indicates blend of slag and fly ash. The curing temperature was 20 °C. The water to precursor ratio was 0.4.



**Figure 14** Conceptual models to describe the concentrations of Al, Ca, K, Fe, Mg, Si, Na, OH<sup>-</sup> and S in the pore solution of alkali-activated slag/fly ash paste. In the graphs,  $[Si]_0$  is the concentration of Si in the alkaline activator.  $[Si]_{max}$  is the maximum concentration of Si.



**Figure 15** Heat evolution rates of alkali-activated fly ash pastes at 40 °C (A) and 60 °C (B). In the graphs,  $P_1$  and  $P_2$  refer to the first and second calorimetric peak respectively, N and S indicate the weight percentage of Na<sub>2</sub>O and SiO<sub>2</sub> with respect to fly ash, and T indicates the curing temperature in Celsius. The water to fly ash ratio was 0.35.

- 0.



Figure 16 The influence of an increase of temperature on the concentrations of Al, Ca, K, Fe and Mg in the pore solution of alkali-activated fly ash paste.  $T_1$  and  $T_2$  are temperatures.



Figure 17 FTIR spectra of alkali-activated fly ash samples at 28 days. In the graph, N and S indicate the weight percentage of  $Na_2O$  and  $SiO_2$  with respect to fly ash, and T indicates the curing temperature in Celsius. The water to fly ash ratio was 0.35.