

Early structural build-up behavior, setting mechanism, and nanostructure of alkali-activated GGBFS mixtures

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3 **Early structural build-up behavior, setting mechanism and**
4 **nanostructure of alkali-activated GGBFS mixtures**

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19 **Abstract**

20 The initial changes in the microstructure of Alkali-activated slag cement (AAS) characterize the
21 rheological behavior, initial structuration and setting times of the mixture. In this study, the
22 relationships between the rheological behavior, solidification process and nanostructure changes of
23 sodium hydroxide-activated slag (NH-AAS) and sodium silicate-activated slag (SS-AAS) pastes over
24 time are investigated by small amplitude oscillation test, isothermal calorimetry, SEM analyses and
25 ¹H-²⁹Si cross-polarization magic angle spinning nuclear magnetic resonance spectroscopy (CP MAS
26 NMR) and ²⁹Si MAS NMR spectroscopy. Test results showed that the NH-AAS reaches its initial
27 setting time following a continuously proceeding structuration process, while the SS-AAS reaches its
28 initial setting with an abrupt increase in the structuration process. At their initial setting times, the NH-
29 AAS and SS-AAS mixtures release similar heat and reach a similar reaction degree. The SS-AAS
30 mixture had more N-A-S-H formation than C-A-S-H at the initial setting time. On the other hand, the
31 amount of C-A-S-H was similar to N-A-S-H in the NH-AAS mixture. The gradual N-A-S-H gel
32 degradation to form C-A-S-H was observed during the acceleration period for the SS-AAS mixture.
33

34 **Keywords:** *Structural build-up, Setting time, Nanostructure, Nuclear magnetic resonance*
35 *spectroscopy (NMR), Alkali-activated cements, Ground granulated blast furnace slag.*
36

37 **1. Introduction**

38 Alkali-activated cement (AAC) has been considered as a low carbon footprint alternative to Portland
39 cement (PC) due to its higher mechanical properties, better volume stability, and greater resistance to
40 sulfate and acid attack depending on the precursor type [1]. Alkali-activated slag cement (AAS) is a
41 common and promising type of AAC, as their production uses ground granulated blast furnace slag
42 (GGBFS), an industrial by-product, and substantially reduces the CO₂ emission [2], [3]. However,
43 although numerous studies have explored the mechanical properties and durability of AAS [2]–[5],
44 very few studies have focused on their microstructural changes over time influencing the rheological
45 properties and setting behavior [6], [7].

46

47 The rise in structural build-up of cementitious materials results from the combination of physical
48 interactions and chemical hydration [7], [8]. In Portland cement, the setting process can roughly be
49 divided into three stages: (i) a colloidal percolation network is formed due to colloidal interactions
50 between cement particles, (ii) cement particles are rigidly connected by the early reaction products
51 such as nano-size C-S-H bridges, (iii) C-S-H bridges are further strengthened, leading to a percolated
52 rigid network [9]. Recently, a few authors have also explained the increase in the early structural
53 build-up of AAC. Alnahhal et al. [10] compared the viscoelastic behavior between alkali-activated
54 slag/fly ash and cement pastes and found that the colloidal attraction between cement particles was
55 significantly stronger than the cohesiveness of the very early inter-particle gel contacts generated in
56 alkali-activated slag/fly ash pastes. They stated that the accumulation of initial reaction products was a
57 possible reason for reaching the solid-like state in the sodium silicate-activated mixtures. In contrast,
58 reaching the solid-like state resulted from a well-percolated network formation in cement paste.
59 Furthermore, another study [8] also pointed out that sodium hydroxide-activated mixtures could also
60 present a percolation network in the early times, a similar setting process with cement pastes. Kashani
61 et al. [11] also investigated the relationship between surface chemistry and rheology in AAS pastes.
62 The authors found that the yield stress was dependent on the changes in the surface chemistry by the
63 amount of activation solutions. For example, a small addition of sodium silicate increased the zeta

64 potential, showing the increase in double-layer repulsive forces reducing the yield stress of AAS
65 pastes. On the other hand, a further increase in sodium silicate amount increased the yield stress,
66 possibly due to the increase in pH causing more slag dissolution resulting in the formation of reaction
67 products.

68

69 The understanding of the change in the nanostructure of AAS from mixing to initial setting time is
70 also crucial. Palacios et al. [12] reported that the rapid loss of fluidity and fast setting of sodium
71 silicate-activated mixtures is due to the initial precipitation of ill-defined N-A-S-H and C-N-A-S-H
72 products. Li et al. [13] used various combined activators to understand the setting and hardening
73 control of AAS. They elucidated that Ca (and hence also Mg) acts as a network modifier in slag glass,
74 causing these precursors to dissolve more quickly. Fly ash, on the other hand, does not exhibit the
75 same level of reactivity due to its chemical stability.

76

77 Acquiring knowledge of the nanostructure of AAS will also enable the industry to develop new
78 strategies to design AAS mixtures and promote their applications. The nanostructure characterization
79 of AAS can be done in different ways. For example, Chen et al. [14] used Fourier-transform infrared
80 spectroscopy (FTIR) and observed that the Q² band shifted to larger wavenumbers and became
81 broader with an increase of silicate modulus in the activator solution. Puertas et al. [15] used various
82 techniques, such as X-ray diffraction (XRD), FTIR and nuclear magnetic resonance (NMR), to
83 determine the structure and composition of the main reaction products. Ismail et al. [16] used
84 environmental scanning electron microscopy (ESEM) with energy dispersive X-ray (EDX) to
85 determine the gel composition. They reported that the silicate-activated slag showed a nanostructure
86 dominated by a C-A-S-H type gel. In contrast, silicate-activated fly ash was dominated by N-A-S-H
87 'geopolymer' gel formations. However, all of the abovementioned studies have focused on the later
88 ages of AAC and cannot provide enough information about the microstructural changes at very early
89 ages.

90 Therefore, this study aims to determine the early gel composition of AAS at the initial setting time and
91 investigates the relationship between the early structural build-up and nanostructure of AAS. The

92 structural build-up of AAS has been evaluated by small amplitude oscillation sweep (SAOS) tests. The
93 nanostructure of AAS was determined by the combined use of ^1H - ^{29}Si cross-polarization magic-angle
94 spinning NMR (CPMAS NMR) and ^{29}Si MAS NMR. The rheological behavior and gel composition of
95 sodium hydroxide-activated (NH-AAS) and sodium silicate-activated slag mixtures (SS-AAS) at the
96 initial setting time was compared.

97

98 **2. Materials and Methods**

99 *2.1 Materials and mix design*

100 In this investigation, ground granulated blast furnace slag (GGBFS) was employed as precursor. This
101 material was provided by ORCEM, the Netherlands. Table 1 displays the results of the chemical
102 composition analyzed by X-ray fluorescence (XRF). The X-ray diffraction pattern and particle size
103 distribution of GGBFS are displayed in Fig. 1. A broad diffraction hump between 20° and 35° 2θ is
104 observed due to the presence of the amorphous components and the absence of crystalline phases.
105 GGBFS has a volume-mean particle sizes (d_{50}) of around $9\ \mu\text{m}$. Sodium hydroxide pellets (NaOH,
106 caustic soda) with 97 % purity and liquid sodium silicate solution were used as activating agents. The
107 liquid sodium silicate solution consists of 28.5% SiO_2 , 17.95% Na_2O and 53.55% water with an Ms
108 value ($\text{SiO}_2/\text{Na}_2\text{O}$) of 1.6. Sodium hydroxide was used to adjust the Ms value of the activator solution
109 to 1.2. The mixture compositions are given in Table 2. Two AAS mixtures were produced in this study;
110 the mixtures coded as NH-AAS was activated by NaOH, while the mixtures coded as SS-AAS was
111 activated by the combination of NaOH and liquid sodium silicate with a Ms value of 1.2. The water-
112 to-solid binder ratio (w/sb) and sodium content ($\text{Na}_2\text{O}\%$) of the mixtures NH-AAS and SS-AAS were
113 0.45 and 5%, respectively. The pH values of activator solutions (Table 2) were determined by a pH
114 meter at $20\ ^\circ\text{C}$. However, it should be noted that using a pH meter to measure the pH of the activator
115 solutions may result in some errors owing to the extremely high alkali ion concentrations and the
116 presence of silica in the activator solution [17].

117

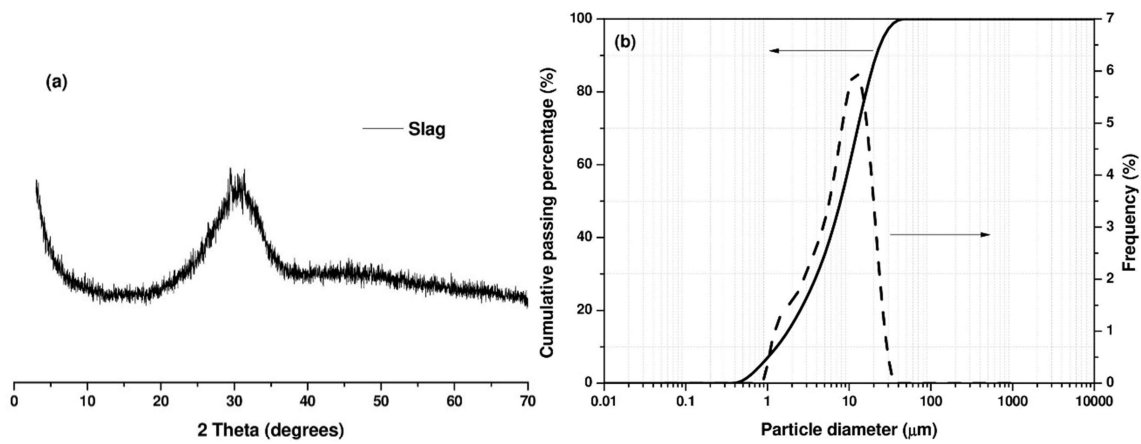
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119

120 Table 1. Chemical composition of the GGBFS.

Precursor	CaO	SiO ₂	Al ₂ O ₃	MgO	SO ₃	TiO ₂	K ₂ O	Na ₂ O	Fe ₂ O ₃	MnO	Others
GGBFS	40.8	33.3	12.3	7.84	2.30	1.29	0.67	0.44	0.39	0.36	0.31

121



122

123 Fig. 1. X-ray diffraction pattern (a) and Particle size distribution (b) of GGBFS.

124

125 Table 2. Mixture design of AAS.

Mixture notation	Activator nature	W/SB ratio	Ms ratio	Na ₂ O (%)	SiO ₂ (%)	pH	Initial setting time (min)
NH-AAS	Sodium hydroxide	0.45	0	5	0	13.82	120
SS-AAS	Sodium hydroxide and sodium silicate	0.45	1.2	5	6	13.50	45

126

127 *2.2 Sample preparation*

128 The activator solutions were prepared one day before usage to ensure the complete dissolution of

129 NaOH. To prepare the paste samples for rheological tests and microstructural analysis, the GGBFS

130 and the activator solution were mixed in a plastic container using a mechanical stirrer for 30 s at 200
131 rpm followed by 3 min at 1800 rpm.

132

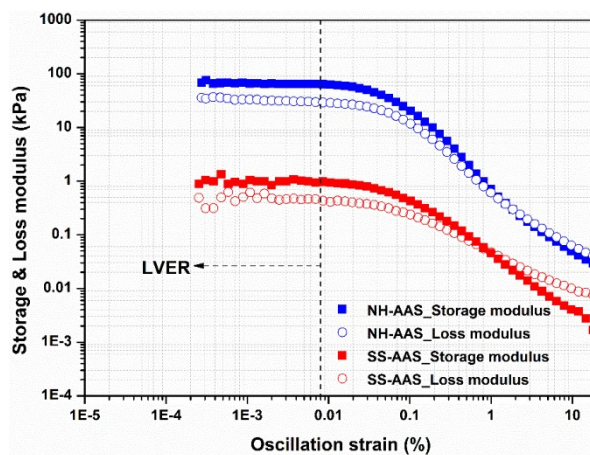
133 2.3 Experimental methods

134 2.3.1 Structural build-up and setting times of AAS pastes

135 Small amplitude oscillation shear (SAOS) tests were carried out in an oscillatory rheometer (Anton
136 Paar MCR 102) to assess the structural build-up of AAS pastes. Storage modulus measurements were
137 used as an indicator of the elastic portion of the mixtures. All the tests in this study were carried out at
138 a temperature of 20 °C.

139 The testing geometry was a six-blade vane in a steel cylindrical cup, which was used to eliminate the
140 effect of slippage between the paste and the plate or the cylindrical cup during measurements [7], [18].
141 A plastic cover was placed on the top of the cup to prevent evaporation. Prior to time-sweep tests, the
142 linear viscoelastic region (LVER) was determined by the strain-sweep test, which was a test that the
143 strain amplitude was increased from 0.0001% to 20% with a constant frequency of 1 Hz. The strain-
144 sweep test results are presented in Fig. 2. The strain amplitude of 0.005% was chosen in time-sweep
145 tests to ensure two mixtures were tested in LVER [6], [7], [19], [20]. The AAS paste was then
146 subjected to pre-shear at 100 s⁻¹ for 30 s and rest for another 30 s to eliminate the residual stress. The
147 SAOS time-sweep tests were immediately started at the age of 6 min.

148



149

150 Fig. 2. Strain-sweep of AAS pastes for the determination of the LVER.

151

152 An automatic Vicat apparatus was used to determine the setting times of AAS pastes. The initial
153 setting times of AAS pastes were determined when the needle penetration depth reached 6 ± 3 mm
154 according to EN196-3 standard [21].

155

156 *2.3.4 Heat evolution of AAS pastes*

157 The calorimetric curves of AAS pastes were measured by eight-channel isothermal calorimetry (TA
158 instrument, USA). After mixing, the AAS pastes were immediately placed into a glass ampoule;
159 afterward, the ampoule was sealed and loaded into the isothermal calorimetry. The calorimetric
160 measurements were carried out at a temperature of 20 °C for 3 days.

161

162 *2.3.5 ^1H - ^{29}Si solid-state cross-polarization magic angle spinning nuclear magnetic resonance 163 (CPMAS NMR) and ^{29}Si MAS NMR analysis*

164 It is important to select an appropriate technique to stop the reactions prior to ^1H - ^{29}Si CPMAS NMR
165 and ^{29}Si MAS NMR tests. In this study, the method to arrest the hydration of AAS was followed by
166 Chen et al. [23]. Firstly, the fresh pastes were mixed with deionized water and then subjected to
167 centrifugation to remove the liquid. This water extraction was repeated two times to remove the
168 soluble silicate species. Afterwards, a chemical extraction was carried out in a way that the solid part
169 was put on the 0.2 μm nylon filter membrane, and then at least 400 ml solvent (50/50 volume of
170 methanol/acetone) was added to wash the solid part and remove the water. Before NMR analysis, the
171 residual particles were collected and stored in a mild vacuum condition for 24 hours. Palacios et al.
172 [12] also noted that the initial washing of the AAS pastes with water and subsequent treatment with
173 isopropanol also can be an appropriate method to achieve the stoppage reactions in AAS pastes.

174

175 ^1H - ^{29}Si solid-state CPMAS NMR and ^{29}Si MAS NMR spectra were acquired on an Agilent VNMRS
176 Direct Drive 400 MHz spectrometer (9.4 Tesla) equipped with a T3HX 3.2 mm probe. The signal of
177 talc for both two types of NMR tests was used to calibrate the silicon chemical shift scale (-98 ppm).
178 Magic angle spinning was performed at 8 kHz and 6.5 kHz using ceramic zirconia rotors for ^1H - ^{29}Si
179 CP MAS NMR and ^{29}Si MAS NMR test, respectively. Acquisition parameters in ^1H - ^{29}Si CP MAS

180 NMR (or ^{29}Si MAS NMR) used were: a spectral width of 24 kHz, a 90° pulse length of 3.2 μs (or 4.5
181 μs), an acquisition time of 15 ms, a recycle delay time of 3 s (or 20 s), a spin-lock field of 80 kHz, a
182 contact time of 1 ms (or a recycle delay time of 20 s), and about 100000 accumulations (or about 8000
183 accumulations). High-power proton dipolar decoupling during the acquisition time for both two types
184 of NMR tests was set to 80 kHz. The spectra of ^1H - ^{29}Si CPMAS NMR and ^{29}Si MAS NMR were
185 decomposed by the Gaussian peaks, and the specified full width at half height was always less than 10
186 ppm [24].

187

188 *2.3.6 Scanning electron microscopy of AAS pastes*

189 SEM with secondary electron mode were used to observe the morphology of AAS pastes. The
190 stoppage method of the reactions was elucidated in the previous section. The specimens at different
191 reaction times were carbon coated before imaging.

192

193 **3. Results and discussions**

194 *3.1 Structural build-up behavior and initial setting times of AAS pastes.*

195 Fig. 3a presents the evolution of the early storage modulus of AAS pastes activated by sodium
196 hydroxide and sodium silicate. As seen in Fig. 3a, these two mixtures exhibited completely different
197 structural build-up behavior. The increase of structural build-up of sodium hydroxide-activated slag
198 paste started off with a greater storage modulus, followed by a steadily rising rate of structural build-
199 up over time. The trend of the structural build-up behavior is quite similar to that of Portland cement.
200 Previous studies have reported that a sharp increase in the storage modulus is observed in the first 30
201 min for Portland cement due to the formation of the percolated network [9], [10], [25]. After that, the
202 storage modulus increases at a relatively lower rate, indicating the sequential development of
203 hydration products at cement particle contact locations [9]. Due to the high alkaline environment for
204 the mixture NH-AAS, the rapid reaction rate of GGBFS could also enhance the formation of reaction
205 products, furthermore inducing the occurrence of the percolated network [6]. In the meantime, the
206 rapid reaction product formation could cover slag's surface, preventing further slag reaction [7].

207 Therefore, a continuous increase in the storage modulus of NH-AAS could be observed. It could then
208 be considered that the setting process of NH-AAS follows a similar mechanism as that of Portland
209 cement [20].

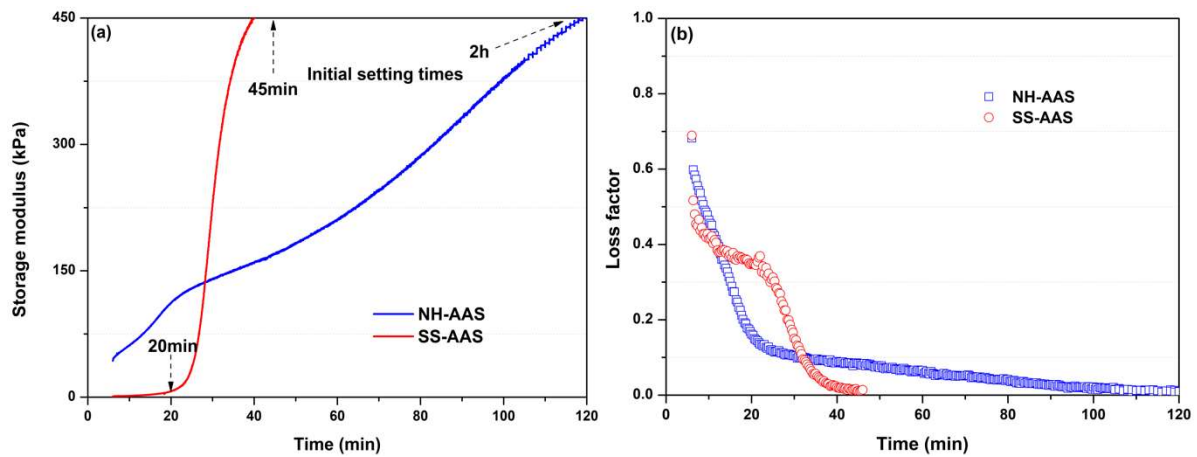
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211 However, the evolution of the structural build-up of SS-AAS was significantly different from the case
212 of NH-AAS. The initial storage modulus started from a very low value as compared to NH-AAS. In
213 the first 20 min, almost no structural formation could be observed for the mixture SS-AAS. However,
214 afterwards, the storage modulus of SS-AAS steeply increased. Some researchers attribute the very low
215 initial storage modulus of SS-AAS to the viscous nature of sodium silicate solution with a great
216 number of aqueous silicates, which prevents the formation of the percolated network [10], [26]. The
217 mild alkaline environment provided by the sodium silicate solution allowed a considerable time to
218 dissolve GGBFS, providing sufficient calcium and aluminum. Once these ions reach their critical
219 concentration, they will interact with the silicates originated from the activator solution, significantly
220 leading to a significantly larger number of reaction products in the matrix [7], [10], [12]. Therefore, a
221 rapid increase in storage modulus could be observed for the SS-AAS mixtures. It should be noted here
222 that the initial setting times of AAS pastes were also consistent with the trend of the structural build-
223 up. The mixture of SS-AAS showed a short initial setting (45 min) as compared to that of NH-AAS
224 (120 min).

225

226 Fig. 3b presents the evolution of loss factor of NH-AAS and SS-AAS mixture. In first 20 min, the loss
227 factor for NH-AAS mixture decreased dramatically, afterwards, the downward trend started to slow
228 down until the loss factor dropped to 0, reflecting that the paste reached solid-like state at around
229 initial setting time (120 min). By contrast, mixture SS-AAS showed an inverse trend. The loss factor
230 for mixture SS-AAS decreased slowly in the first 20 min, indicating its more viscous behavior. Then,
231 a rapid decrease could be observed for SS-AAS mixture until its initial setting time (45 min). This
232 showed that SS-AAS mixture reached solid-like state earlier than NH-AAS mixture.

233



234

235 Fig. 3. Evolution of storage modulus (a) and loss factor (b) of sodium hydroxide-activated and sodium silicate-
 236 activated GGBFS pastes.

237

238 *3.2 Calorimetric investigation of AAS pastes.*

239 The evolution of calorimetric curves, including heat flow and cumulative heat release of AAS pastes,

240 is presented in Fig. 4. As well known, there are five distinct stages for the AAS mixtures: dissolution,

241 induction, acceleration, deceleration and steady-state [29]–[32]. As demonstrated in Fig. 4a, the AAS

242 mixtures exhibited a high local peak during the first 20 min. Rapid heat evolution was observed when

243 the activator solution was in contact with GGBFS, being mostly related to the wetting and dissolution

244 of GGBFS particles and partly related to the interaction of silicate units and sodium with calcium and

245 aluminum dissolved from GGBFS [33], [34]. For SS-AAS pastes, the dissolution stage was followed

246 by an induction period in the heat flow, while no such period was seen for the NH-AAS pastes. The

247 observed acceleration peak (second peak) after the induction period for the AAS pastes is usually

248 associated with the onset of the main strength and microstructural development [35]. The occurrence

249 of the acceleration peak of SS-AAS was delayed as compared to the NH-AAS mixture, and the

250 intensity of the acceleration peak in the SS-AAS was lower as compared to the NH-AAS mixture,

251 which is also consistent with previous studies [29], [33]. This could be related to the lower pH value of

252 the sodium-silicate activator solution compared to the sodium-hydroxide activator (Table 2). It was

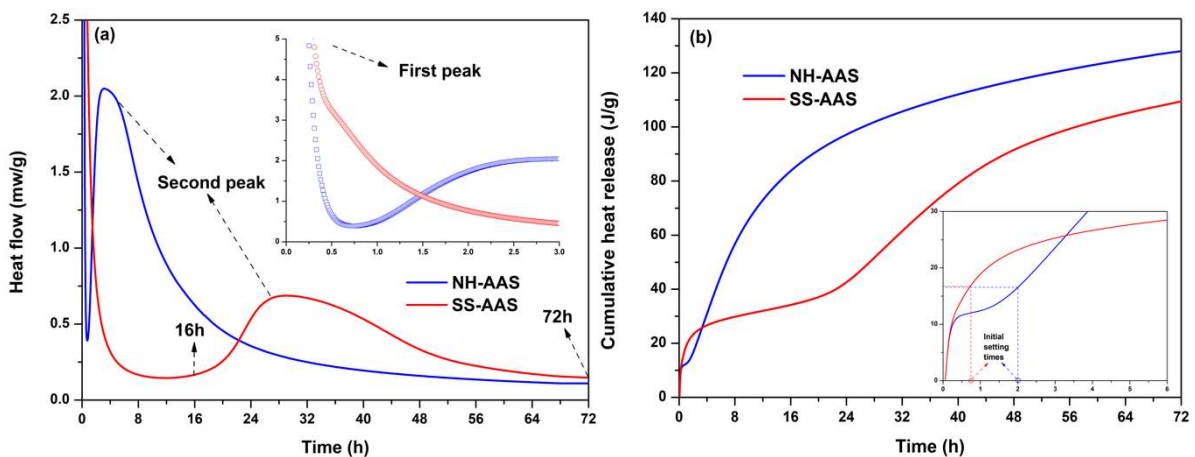
253 reported that the higher alkaline environment normally enhances the dissolution rate and slag reaction

254 degree [36]. Additionally, increasing alkalinity also improves the solubility of silica and alumina in

255 solution, which could induce more reaction product formation [27], [37].

256

257 The cumulative heat release of NH-AAS showed a higher value as compared to that of SS-AAS at the
258 age of 72 h. However, in the first 3.5 h, the SS-AAS released more heat than NH-AAS. This is
259 probably due to the interaction between the calcium ions, continuously released from GGBFS, and the
260 silicate ions, originating from sodium silicate solution, quickly forming large amounts of gel products
261 [38]. This is also consistent with the rapid increase in storage of SS-AAS in the early time. On the
262 other hand, NH-AAS and SS-AAS both presented a similar cumulative heat release of around 17 J/g at
263 their initial setting times (Fig. 4b).



264

265 Fig. 4. Evolution of heat flow (a) and cumulative heat release (b) of sodium hydroxide-activated and sodium
266 silicate-activated GGBFS pastes.

267

268 3.3 NMR analysis of AAS pastes

269 To further understand the relationships between the nanostructure and the setting process of AAS,
270 NMR tests were carried out on the samples taken at critical reaction times based on storage modulus,
271 setting time and calorimetric measurements. Three samples were taken at 20 min and 45 min (initial
272 setting time, as shown in Table 2) for the SS-AAS mixture and at 120 min (initial setting time) for the
273 NH-AAS mixture based on storage modulus and setting times as shown in Fig. 3. In addition, two
274 samples were taken from the SS-AAS mixture corresponding to the beginning of the acceleration stage
275 (16 h) and the end of the deceleration period (72 h), based on the calorimetry measurements as shown
276 in Fig. 4.

277

278 3.3.1 NMR analysis at initial setting times

279 The ^{29}Si MAS NMR spectra of the anhydrous GGBFS and the pastes at different reaction times are
280 given in Fig. 5. As shown in Fig. 5, the anhydrous slag was the primary source of the signal at -76.8
281 ppm in all the spectra. In addition, a shoulder between -80 and -105 ppm was observed at the initial
282 setting time for NH-AAS (120 min) and SS-AAS mixtures (45 min), reflecting the reaction product
283 formation. Therefore, these reaction products should be responsible for the initial setting of AAS. No
284 difference was observed between the spectra of SS-AAS_20min and anhydrous GGBFS, indicating no
285 reaction product formation at 20 min. This was also consistent with the results of the structural build-
286 up of the SS-AAS mixture, showing no significant structural formations at 20 min.

287

288 To better understand the nanostructure of AAS, an accurate deconvolution should be performed on the
289 shoulder region. However, since some $\text{Q}^n(\text{mAl})$ ($n=0,1,2,3,4$ and $m=0,1,2,3,4$) peaks were overlapped
290 in the ^{29}Si MAS NMR spectra [39], a ^1H - ^{29}Si CPMAS NMR test was performed to find the peak
291 positions, prior to the deconvolution of ^{29}Si MAS NMR spectra. It should be noted that the chemical
292 shift of each resonance is required to be consistent in both the ^{29}Si MAS NMR and ^1H - ^{29}Si CPMAS
293 NMR spectral deconvolutions.

294

295 Fig. 6 and Fig. 7 present the differences between the ^1H - ^{29}Si CPMAS NMR and ^{29}Si MAS NMR
296 spectra of the pastes activated by sodium hydroxide and sodium silicate, respectively, as well as their
297 detailed deconvolution. The resonances observed in the ^1H - ^{29}Si CPMAS NMR are influenced by their
298 proximity to protons and the Hartmann-Hahn contact period, and so this spectrum particularly shows
299 those Si species present in the hydration reaction products [40]. As a result, the ^1H - ^{29}Si CPMAS NMR
300 data in Fig. 6 and 7 can confirm that the broad resonance feature observed in the ^{29}Si MAS NMR is
301 highly due to the presence of the anhydrous slag in the AAS. It should be noted that although the ^1H -
302 ^{29}Si CPMAS NMR can provide better resolution of the spectra in terms of reaction products, it can
303 only give qualitative information rather than quantitative information. Therefore, the combined use of
304 ^1H - ^{29}Si CPMAS NMR and ^{29}Si MAS NMR enables a more complete examination of the variations in
305 the nanostructure of reaction products that are formed in the AAS mixtures.

306
307 The nanostructure of AAS can be identified by the $Q^n(mAl)$ structural units, showing various chemical
308 environments of silicon tetrahedral of phases in ^{29}Si NMR spectra, where n denotes the number of
309 oxygen bridges between each silicon tetrahedral unit and other silicon atoms, and m represents the
310 number of aluminum tetrahedral occupying bridging positions [41], [42]. The deconvolution results of
311 the $Q^n(mAl)$ structural units in the ^{29}Si MAS NMR spectra are summarized in Table 3. The
312 deconvoluted peak located around -76.8 ppm for NS-AAS and SS-AAS mixtures at initial setting
313 times refers to the remnant GGBFS (Q^0), and the integration of these peaks reached 80.04% and 81.28%
314 of the spectra of ^{29}Si MAS NMR, respectively. In other words, when both NH-AAS and SS-AAS
315 mixtures show an initial setting, they both have a similar reaction degree. Therefore, it may indicate
316 that the initial setting of AAS would occur when the reaction degree of GGBFS approached 20%.
317 Combining the results of rheology and setting time, it can be concluded that the solidification of NH-
318 AAS is a continuous process; however, the initial set of SS-AAS occurs suddenly in a shorter period
319 [20].

320
321 As shown in Fig. 6b, the peak position for the mixture NH-AAS_120 min with $\delta = -79.2$ is attributed
322 to the Q^1 site, which is a terminal silicate tetrahedral site inside C-A-S-H gels [24], [43]. The middle-
323 of-chain silicate inside C-A-S-H gels is connected to the $Q^2(1Al)$ and Q^2 sites at -83.5 ppm and -86
324 ppm, respectively [41], [44]. Since there is an overlap between resonances in this region, the peak at -
325 89.2 ppm belongs to $Q^3(1Al)$ and $Q^4(4Al)$ sites [16], [24], [45]. Herein, it should be noted that the
326 $Q^3(1Al)$ site denoted a substantial degree of cross-linking with the C-(N-)A-S-H gel [24], [40], while
327 $Q^4(4Al)$ site refers to the Al-rich N-A-S-H gel [46]. Additionally, $Q^4(2Al)$, $Q^4(1Al)$, and $Q^4(0Al)$ each
328 have their own set of three extra sites located somewhere in the range of -96 to -110 ppm. The $Q^4(mAl)$
329 environments are attributed to a polymerized Si-rich N-A-S-H gel [12], [40], [47].

330
331 Similarly, as shown in Fig. 7b, three resonances with $\delta = -78.5$ ppm, -83.0 ppm and -86 ppm, are
332 attributed to Q^1 , $Q^2(1Al)$ and Q^2 sites, respectively, indicating the formation of C-A-S-H in the SS-
333 AAS mixture. The $Q^3(1Al)$ and $Q^4(4Al)$ sites are located at -88.6 ppm, showing the formation of high

334 crosslinking C-(N-)A-S-H and N-A-S-H, respectively. In addition, the alumino-silicate gel N-A-S-H
335 can be confirmed by three additional peaks between -93 and -106 ppm, indicating the $Q^4(3Al)$, $Q^4(2Al)$
336 and $Q^4(1Al)$ sites, respectively.

337

338 As shown in Table 3, it is found that the SS-AAS mixture showed a higher percentage of N-A-S-H
339 formation ($Q^4(mAl)$) as compared to C-A-S-H (Q^1 , $Q^2(1Al)$ and Q^2), while the amount of C-A-S-H
340 (Q^1 , $Q^2(1Al)$ and Q^2) was more than N-A-S-H ($Q^4(mAl)$) in the NH-AAS mixture. Although it was
341 very difficult to distinguish the net amounts of $Q^3(1Al)$ and $Q^4(4Al)$ due to their overlapped resonance,
342 it could still be concluded that the SS-AAS mixture contained a higher percentage of N-A-S-H as
343 compared to the NH-AAS mixture at initial setting times based on the other Q species. In addition, it is
344 well known that the sodium silicate solution contains high Q species [13], [17]. Due to the condensed
345 silicates in the activator of SS-AAS, these higher-Q Si species are more likely to condense with Al and
346 Ca to form 3D Ca-modified gel at early ages [48], [49]. While regarding the NaOH-activated mixture,
347 congruent dissolution of the various Si coordination environments in the precursors is highly unlikely
348 to occur during alkali activation due to the presence of many phases in the precursors and the
349 previously observed preference of Al over Si dissolution in the silicoaluminate precursors.

350 Furthermore, the highly polymerized aluminosilicate phase is predicted to disintegrate more slowly
351 than the depolymerized calcium silicate phases observed in the precursors [46], [50]. And these
352 released depolymerized silicates interacted with aluminum and calcium ions to form chain-like C-A-S-
353 H gel [15], [37]. This could be the reason for the distinctive difference between SS-AAS and NH-AAS
354 in terms of gel composition at early ages.

355

356 It is plausible to see that the hydration products of AAS at initial setting times are composed of the C-
357 A-S-H and N-A-S-H gels. In the meantime, the coexistence of C-A-S-H and N-A-S-H resulted in a 3D
358 Ca-modified gel (N,C)-A-S-H, which was also reported by Garcia-Lodeiro et al. [51]. However, very
359 few studies have focused on the gel composition of AAS at the initial setting time in the literature.

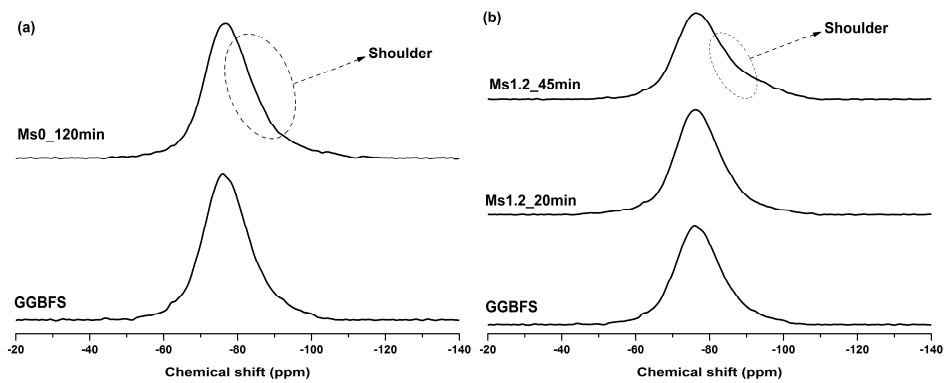
360 Only Palacios et al. [12] used the thermodynamic modeling and NMR technique to confirm the
361 presence of the coexisting formation of the alumino-silicate gel and C-A-S-H gel in the sodium

362 silicate-activated slag mixtures at early times. These C-A-S-H and N-A-S-H gels are formed from the
363 reactions of Ca^{2+} ions and aluminate species dissolved from slag and Na^+ and silicate species initially
364 existed in the solution. Therefore, the rheological properties and the setting process are determined by
365 these reaction products.

366

367 However, many studies, in general, pointed out that the main reaction products of AAS at later ages
368 are C-A-S-H gels. It is, therefore, very worthwhile to investigate when these N-A-S-H gels start to
369 disappear or whether they are converted into C-A-S-H gels as the reaction continues. Therefore, to
370 investigate a possible change in the gel composition of AAS at an early time, NMR tests were
371 conducted on the samples taken at two critical reaction times based on the isothermal calorimetric
372 curves of SS-AAS pastes (as shown in section 3.4.2).

373



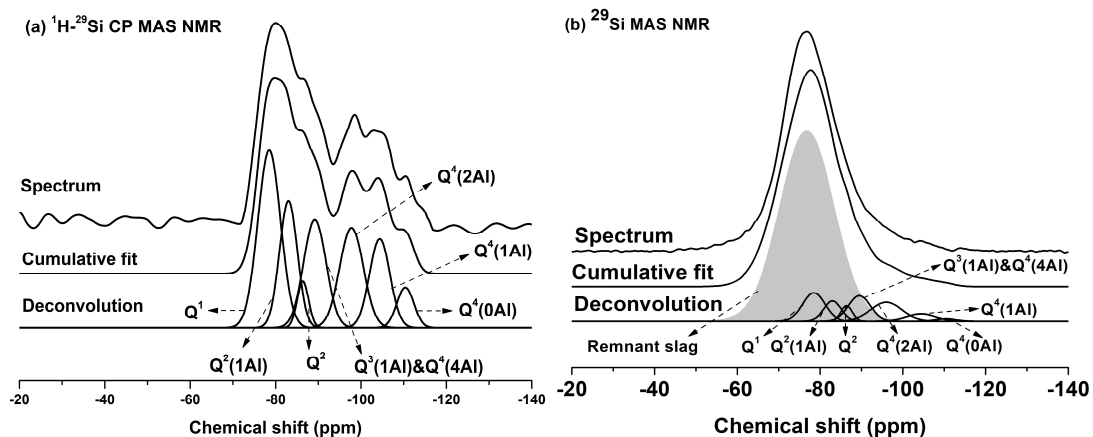
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375 Fig. 5. The original ^{29}Si MAS NMR spectra of AAS, (a) NH-AAS, (b) SS-AAS.

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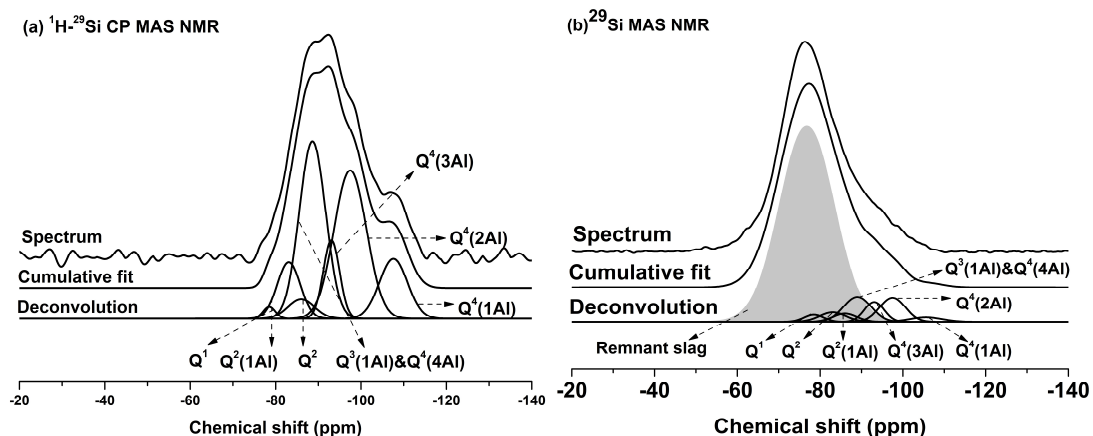
380

Fig. 6. The deconvolution of ^1H - ^{29}Si CPMAS NMR and ^{29}Si MAS NMR spectra of NH-AAS at initial setting

381

time (120 min), (a) The deconvolution of ^1H - ^{29}Si CPMAS NMR, (b) The deconvolution of ^{29}Si MAS NMR.

382



383

384

385

Fig. 7. The deconvolution of ^1H - ^{29}Si CPMAS NMR and ^{29}Si MAS NMR spectra of SS-AAS at the initial setting

386

time (45 min), (a) The deconvolution of ^1H - ^{29}Si CPMAS NMR, (b) The deconvolution of ^{29}Si MAS NMR.

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395 Table 3. The deconvolution results of the ²⁹Si MAS NMR spectra of AAS mixtures over time based on area
 396 percentage (%).

Mixtures	Unreacted		Reaction Products						
	slag		C-A-S-H/C-(N-)A-S-H				N-A-S-H		
	Q ⁰	Q ¹	Q ² (1Al)	Q ²	Q ³ (1Al)	Q ⁴ (3Al)	Q ⁴ (2Al)	Q ⁴ (1Al)	Q ⁴ (0Al)
NH-	80.04	4.93	2.74	1.34	4.38	-	4.59	1.62	0.38
AAS_120min	(-76.8 ppm)	(-78.5 ppm)	(-83.0 ppm)	(-86.4 ppm)	(-89.3 ppm)		(-96.5 ppm)	(-104.4 ppm)	(-110.4 ppm)
SS-	81.28	1.07	1.93	1.51	5.32	2.99	4.75	1.14	-
AAS_45min	(-76.8 ppm)	(-78.5 ppm)	(-83.1 ppm)	(-86.0 ppm)	(-89.0 ppm)	(-93.1 ppm)	(-97.2 ppm)	(-105.6 ppm)	
SS-AAS_16h	68.68	2.45	3.80	4.65	7.63	5.34	4.85	2.60	-
	(-76.8 ppm)	(-78.5 ppm)	(-83.1 ppm)	(-86 ppm)	(-88.9 ppm)	(-93.1 ppm)	(-97.3 ppm)	(-105.6 ppm)	
SS-AAS_72h	48.46	18.22	13.58	8.09	5.55	3.68	2.42	-	-
	(-76.8 ppm)	(-78.8 ppm)	(-83.6 ppm)	(-86.2 ppm)	(-89.1 ppm)	(-93.6 ppm)	(-97.5 ppm)		

397

398

399 *3.3.2 NMR analysis of SS-AAS pastes based on calorimetric curves*

400 Fig. 8 illustrates the ²⁹Si MAS NMR of SS-AAS pastes over time. As the curing age increased from
 401 the beginning to 16 h, the broad peak did not shift significantly, while the shoulder area between -80
 402 and -105 ppm was enhanced. With time elapsing from 16 h to 72 h, the broad peak started to shift to a
 403 higher value, and the shoulder area between -90 and 105 ppm significantly decreased. This is possibly
 404 due to the increase of polymerization degree and cross-linking of gels over time and the disappearance
 405 of the alumino-silicate gel.

406

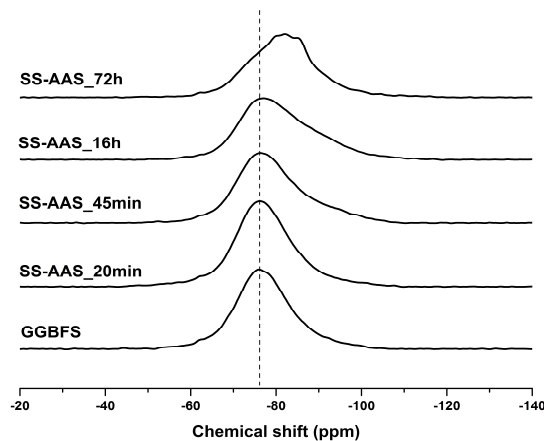
407 ²⁹Si MAS NMR decomposition results of the SS-AAS mixture at around the starting time of the
408 acceleration peak and the end of the deceleration peak observed in the isothermal calorimetric curves
409 are presented in Fig. 9. The area percentage of various Qⁿ(mAl) structural units are quantified and
410 summarized in Table 3. As can be seen from Fig. 9, Q²(1Al) and Q² sites within C-A-S-H increased
411 from 3.80 % to 13.58 % and from 4.65 % to 8.09 % with increasing curing time from 16 h to 72 h,
412 respectively. In particular, the content of Q¹ significantly increased from 2.45 % to 18.22 % with time
413 elapsing and became the most dominant Qⁿ(mAl) structural unit in SS-AAS pastes at the age of 72 h.
414 This is also consistent with the previous studies, reporting the dissolved calcium and aluminum from
415 slag enhance the gel polymerization and result in more crosslinked C-A-S-H gels [24]. During the
416 proceeded reaction time from the initial setting time (45 min) to 16 h and then to 72 h, the content of
417 Q³(1Al) & Q⁴(4Al) gradually increased from 5.32 % to 7.63 % and then decreased to 5.55 %.
418 Meanwhile, the content of N-A-S-H gel (including Q⁴(mAl), m=3, 2, 1, 0) first increased from 8.88 %
419 to 12.79 % when the age increased from 45 min to 16 h and then decreased to 6.10 % when the age
420 increased from 16 h to 72 h. This indicates that the system is continuously enriched with N-A-S-H
421 until the end of the induction period; however, as the reaction proceeded, the N-A-S-H started to
422 destabilize to form C-A-S-H when reaching the end of the second exothermic peak of SS-AAS.
423

424 Based on the NMR results in this study and the compatibility between C-A-S-H and N-A-S-H gels
425 [51], the activation process of SS-AAS pastes in the early times can be explained as follows. After
426 mixing the GGBFS with sodium silicate solution, the process begins with the dissolution of the
427 GGBFS particles in the alkaline solution via rupture of the T-O-T bonds (T: Si or Al) and Ca-O bonds
428 in the slag, resulting in a wide variety of dissolved species. This can be confirmed by various previous
429 studies, which use ICP-OES to observe the change of different ion concentrations [6], [7], [13], [15],
430 [27], [37]. When the ions reach their oversaturation, a great number of C-A-S-H and N-A-S-H gels
431 precipitate. These gels generally contain a high aluminum content, as shown in the NMR results in this
432 study. As the alkaline reaction proceeds, the silicates from the original activator solution are consumed
433 during the formation of reaction products. Therefore, the Si-O bonds in GGBFS start to dissolve and
434 keep the silicon concentration at a high level. In this study, from the initial setting time to 16 h, Si

435 uptake enhances the gel condensation in terms of C-A-S-H, leading to an increase in $Q^2(1Al)$. In the
436 meantime, the percentage of N-A-S-H gel also increases when the age increases from the initial setting
437 time to 16 h.

438

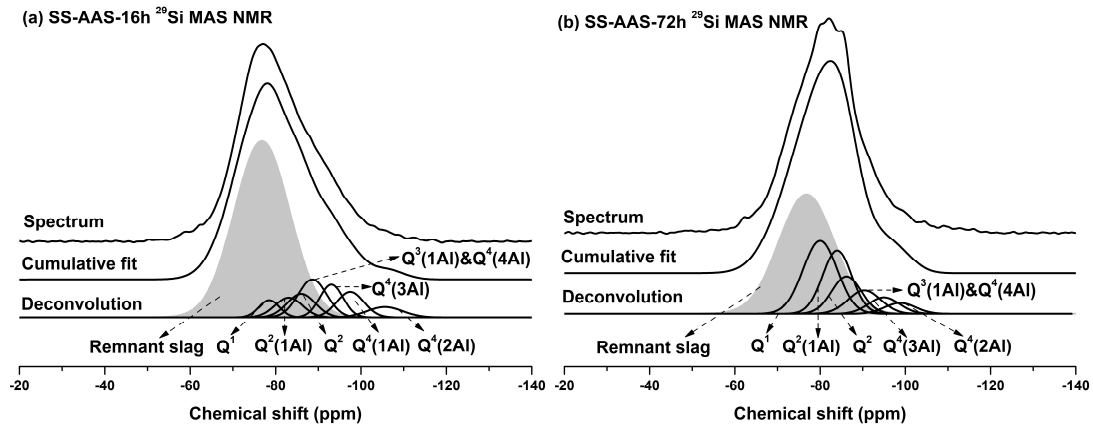
439 Additionally, with time elapsing, the calcium and aluminum ions in the aqueous solution begin to
440 diffuse throughout the cementitious matrix and a certain level of Ca ions reacts with N-A-S-H to form
441 (N,C)-A-S-H. Since calcium and sodium ions have the same ionic radius and electronegative potential,
442 the 3D structure of the (N,C)-A-S-H type gel is maintained by an ion exchange process, in which
443 calcium can replace the sodium ions in a mechanism similar to that observed in clay and zeolites [52].
444 When calcium concentrations are high enough, they diffuse into pores in the matrix and interact with
445 the (N,C)-A-S-H gel. Because of the Ca^{2+} polarising effect, the Si-O-Al bonds are under tension and
446 eventually broken, forming Si-O-Ca bonds. As aluminum is released from the N-A-S-H gel, fewer
447 polymerized structures, such as C-A-S-H gels, are formed. Simultaneously, the C-A-S-H gel formed in
448 earlier phases is capable of absorbing a significant amount of silicon and aluminum ions in bridge
449 positions [53]. Besides, Puertas et al. [15] also found that the calcium concentration and pH value of
450 the waterglass-activated slag mixtures reached a peak value at the age of 24h. This also supports that
451 the N-A-S-H gel will degrade to C-A-S-H due to its low stability in the presence of high pH and high
452 calcium concentrations, as reported by Garcia-Lodeiro et al. [51]. Eventually, the chain-like C-A-S-H
453 gels will form and dominate the reaction products, which is also consistent with this study where Q^1
454 Q^2 and $Q^2(1Al)$ are the main Q species at the age of 72h.



455

456

Fig. 8. The ^{29}Si MAS NMR spectra of anhydrous GGBFS and SS-AAS pastes at different times.



457

458

459 Fig. 9. The deconvolution of ^{29}Si MAS NMR spectra of SS-AAS at around: (a) starting point of the acceleration
 460 period (16 h) and (b) at the end of the deceleration period (72 h) based on isothermal calorimetric curves.

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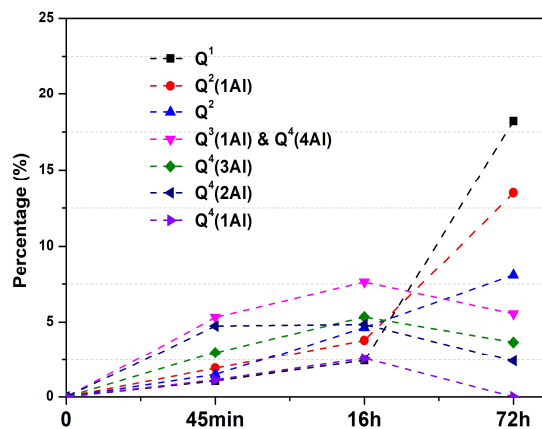
462 3.3.3 Quantitative assessment of C-A-S-H/C-(N-)A-S-H and N-A-S-H

463 Fig. 10 presents the deconvolution results for the ^{29}Si MAS NMR spectra of the SS-AAS mixture over time.

464 It could be seen that in the first 16 h, the $\text{Q}^4(\text{mAl})$ is dominated as compared to other Q^n environments. As curing
 465 time proceeded, the $\text{Q}^4(\text{mAl})$ started to decrease, in particular for the $\text{Q}^4(1\text{Al})$, which decreased to zero at 72 h.

466 However, other chain-like Q^n increased.

467



468

469 Fig. 10. The deconvolution results for the ^{29}Si MAS NMR spectra of SS-AAS mixture over time.

470

471 Richardson and Groves [54] have proposed a conceptual model named “substituted general model” (SGM) to
 472 describe the C-(N-)A-S-H product, which is a mixture of 14 Å tobermorite, jennite, and $\text{Ca}(\text{OH})_2$ structures.

473 However, in recent studies, more formation of cross-linked Si sites, such as Q^3 and $\text{Q}^3(1\text{Al})$, is identified by the

474 high-resolution ^{29}Si MAS NMR [24], [36], [39], [41]. As reported by Myers et al. [24], in tobermorite structural
475 models, Q^3 -type silica bonding environments can only be described by cross-linking between bridging sites in
476 the silicate chains. Therefore, another conceptual model, the “Cross-linked substituted tobermorite model”
477 (CSTM), has been proposed to support the description of the C-(N-)A-S-H gel as a mixture of cross-linked (11 Å
478 tobermorite) and non-cross-linked (14 and/or 9 Å tobermorite) chains. And it should be noted that there are some
479 structural constraints included in the CSTM model: (i) two times as many silicate species exist in the form ($\text{Q}^2 +$
480 $\text{Q}^2(1\text{Al})$) as in the form ($\text{Q}^3 + \text{Q}^3(1\text{Al}) + \text{Al}[4]$). (ii) since aluminum is only substituted into bridging sites in
481 cross-linked tobermorite, the fraction of aluminum substitution into Q^3 type sites is equal to the ratio of $\text{Q}^2(1\text{Al})$
482 to Q^2 sites, and (iii) the introduction of one $\text{Al}[4]$ species into cross-linked tobermorite introduces one $\text{Q}^3(1\text{Al})$
483 and two $\text{Q}^2(1\text{Al})$ Si species. Regarding the structure of N-A-S-H, unfortunately, no acknowledged conceptual
484 model has been proposed due to its complicated 3D structures.

485

486 To further investigate the structure of the investigated samples in this study, two extreme scenarios are
487 considered. In this study, as the $\text{Q}^3(1\text{Al})$ and $\text{Q}^4(4\text{Al})$ are overlapped at around -89 ppm, it was assumed as a first
488 case that all the signals came from $\text{Q}^4(4\text{Al})$ and there was no $\text{Q}^3(1\text{Al})$ in the nanostructure, which means all C-
489 (N-)A-S-H structures consist of non-cross-linked gel as defined by the SGM model. In the second extreme case,
490 it was assumed that the C-(N-)A-S-H had the maximum crosslinking degree (maximum content of $\text{Q}^3(1\text{Al})$, and
491 the remaining part is $\text{Q}^4(4\text{Al})$), considering the abovementioned structural constraints. This indicates that the C-
492 (N-)A-S-H is a mixture of cross-linked and non-cross-linked tobermorite-like structures as defined by CSTM.

493

494 Table 4 presents the C-A-S-H/C-(N-)A-S-H and N-A-S-H gel amounts considering the two extreme scenarios
495 above. As shown in Table 4, the calculated amounts of C-A-S-H/C-(N-)A-S-H and N-A-S-H vary in a narrow
496 band for these two extreme scenarios except the mixture SS-AAS_72h. For example, NH-AAS_120 min
497 exhibited a range of C-A-S-H/C-(N-)A-S-H between 9.01 and 9.68, and N-A-S-H was between 10.30 and 10.97.
498 In both scenarios, the relative amount of C-A-S-H/C-(N-)A-S-H is closer to N-A-S-H gel for the mixture of NH-
499 AAS_120 min, while the N-A-S-H gel is much more dominant in the mixture of SS-AAS_45 min. With time
500 elapsing, the C-A-S-H/C-(N-)A-S-H amount increased rapidly, while the amount of N-A-S-H firstly increased
501 until the end of the induction period based on calorimetric curves. At the end of deceleration period based on
502 calorimetric curves (72h), the N-A-S-H formation for mixture SS-AAS decreased significantly for both two
503 extreme scenarios as compared to the mixture at the age of 16h.

504 Table 4 Nanostructure of investigated pastes in this study considering the two extreme scenarios.

Mixture	Q ¹	Q ² (1Al)	Q ²	Q ³ (1Al)	Q ⁴ (4Al)	Q ⁴ (3Al)	Q ⁴ (2Al)	Q ⁴ (1Al)	Q ⁴ (0Al)	C-A-S- H/C-(N-)A-S-H	N-A- S-H	
No cross-link in	NH-											
C-A-S-H/C-(N)- A-S-H	AAS_120min	4.93	2.74	1.34	-	4.38	-	4.59	1.62	0.38	9.01	10.97
	SS-											
	AAS_45min	1.07	1.93	1.51	-	5.32	2.99	4.75	1.14	-	4.51	14.20
	SS-AAS_16h	2.45	3.80	4.65	-	7.63	5.34	4.85	2.60	-	10.90	20.42
	SS-AAS_72h	18.22	13.58	8.08	-	5.55	3.68	2.42	-	-	39.88	11.65
Maximum cross- link in C-A-S- H/C-(N)-A-S-H	NH-											
	AAS_120min	4.93	2.74	1.34	0.67	3.71	-	4.59	1.62	0.38	9.68	10.30
	SS-											
	AAS_45min	1.07	1.93	1.51	0.27	5.05	2.99	4.75	1.14	-	4.78	13.93
	SS-AAS_16h	2.45	3.80	4.65	0.61	7.02	5.34	4.85	2.60	-	11.51	19.81
	SS-AAS_72h	18.22	13.58	8.08	4.04	1.51	3.68	2.42	-	-	43.92	7.61

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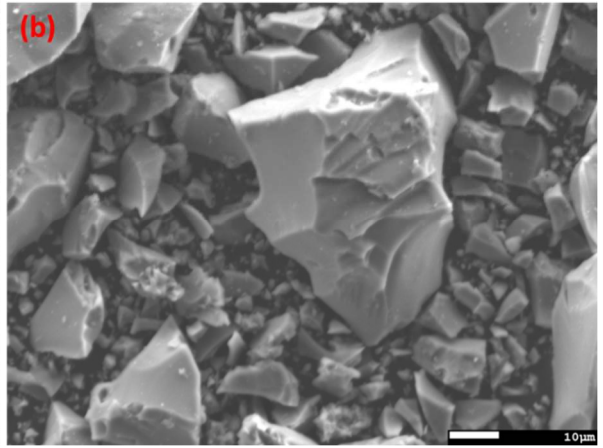
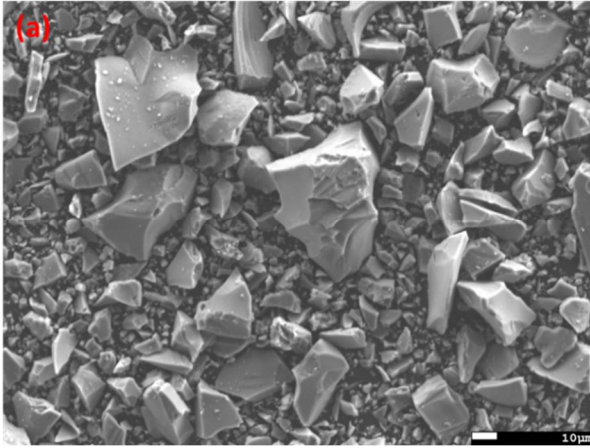
506 *3.4 SEM images of AAS pastes*

507 Scanning electron microscopy (SEM) was used to evaluate the surface morphology development of
 508 the AAS pastes [12], [16]. Fig. 13 presents SEM images of AAS pastes during the reaction process. At
 509 the age of 120 min for NH-AAS (Fig. 13c and 13d) and 45 min for SS-AAS (Fig. 13g and 13h), the
 510 GGBFS particle surfaces became rough due to the precipitation of reaction products, thereby probably
 511 leading to the initial set of AAS pastes.

512

513 As shown in Fig. 13e and 13f, the surface of the GGBFS particles of the SS-AAS mixture at the age of
 514 20 min was very smooth, indicating no detectable reaction product formation at this age. This is also
 515 consistent with the structural build-up and NMR results in this study. With time elapsing, the
 516 dissolved calcium and aluminum ions from the GGBFS reached oversaturation and interacted with the
 517 silicates originating from the activator solution, forming more reaction products. Therefore, the surface
 518 of GGBFS particles became increasingly rough, and more particles would connect to each other due to
 519 the formation of reaction products (as shown in Fig. 13k and 13l) [12].

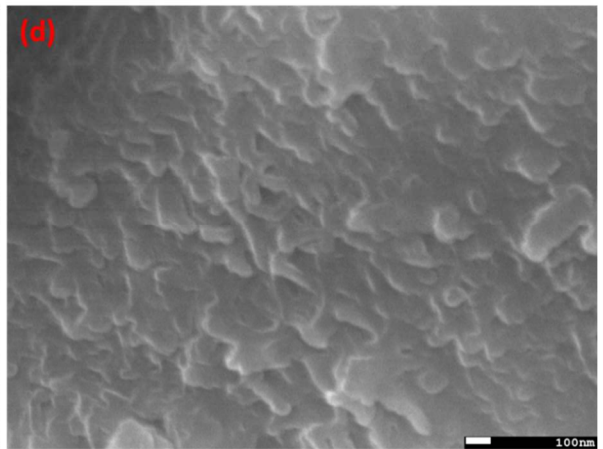
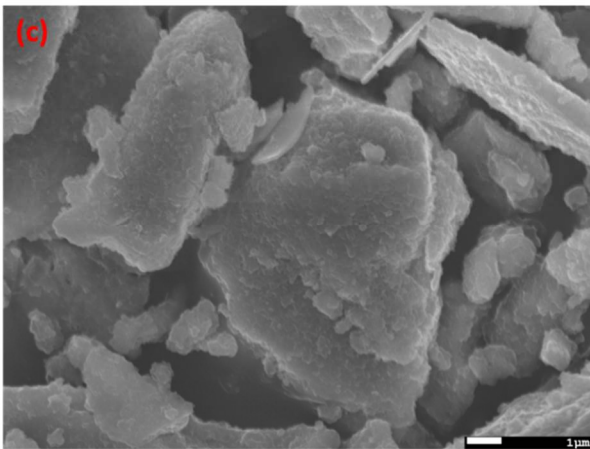
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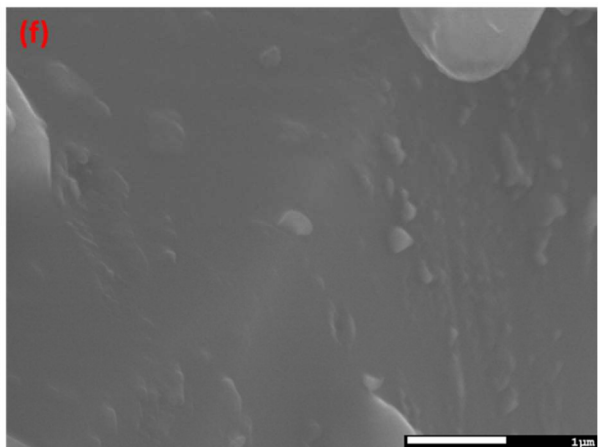
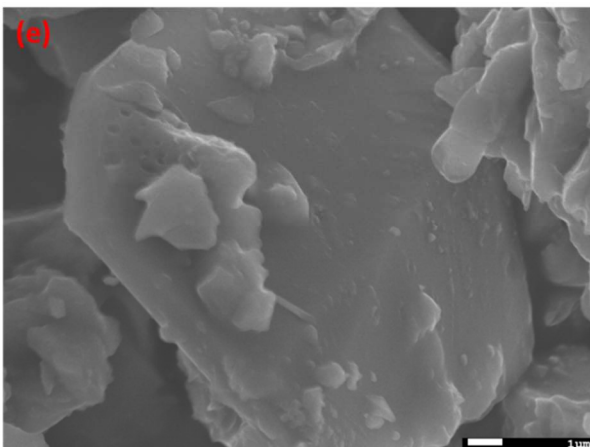
Original GGBFS



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NH-AAS_120min

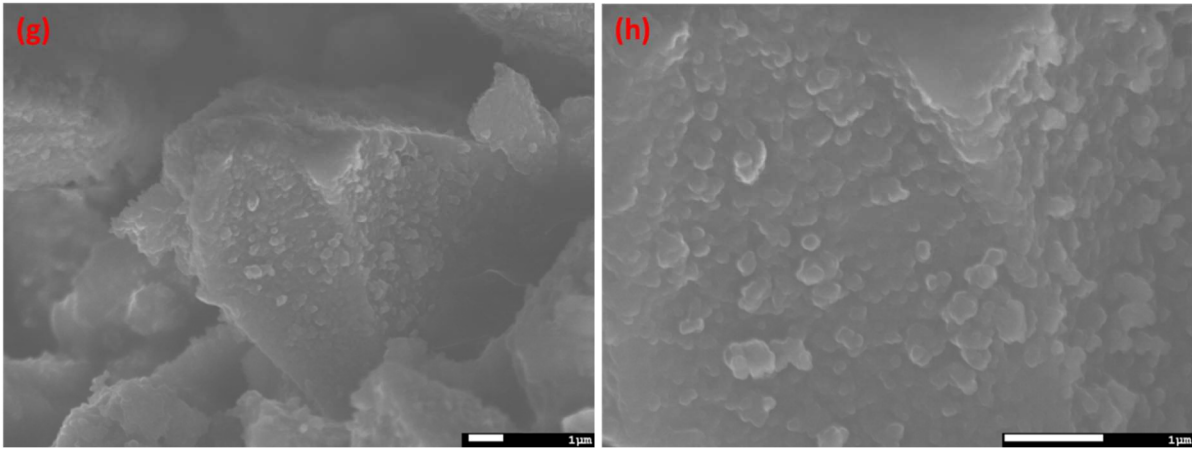


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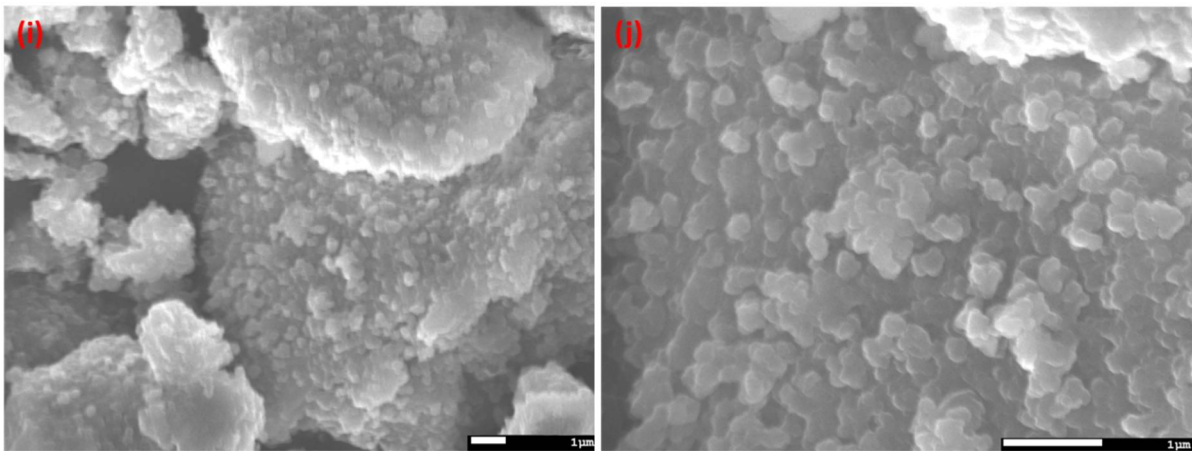
SS-AAS_20min



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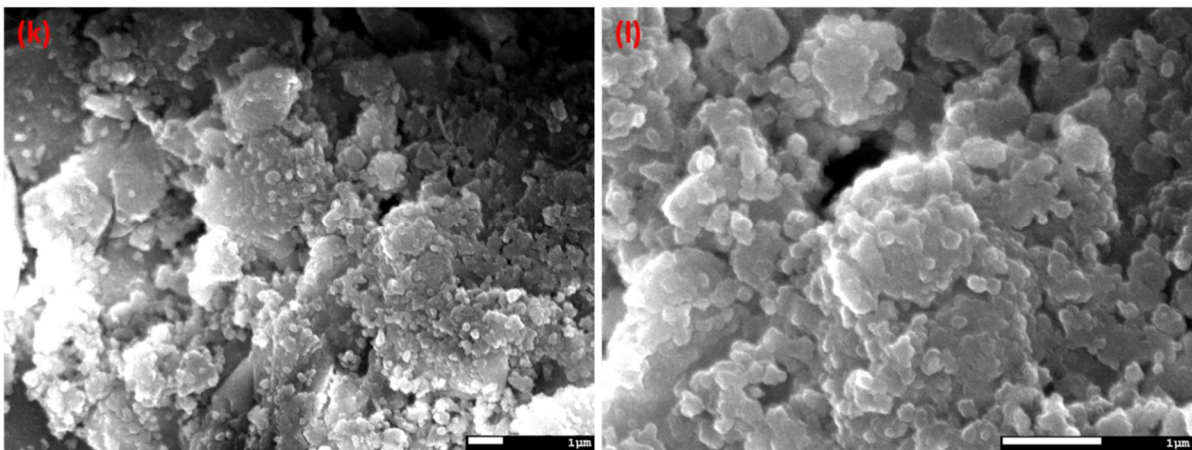
SS-AAS_45min



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SS-AAS_16h



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SS-AAS_72h

534 Fig. 13. SEM images of AAS pastes over time; (a, b) anhydrous GGBFS; (c, d) NH-AAS at 120 min; (e, f) SS-

535

AAS at 20 min; (g, h) SS-AAS at 45 min; (i, j) SS-AAS at 16 h; (k, l) SS-AAS at 72 h.

536

537

538 **4. Conclusions**

539 Following are the primary conclusions that can be derived from the experimental findings:

- 540 • The NH-AAS reaches its initial setting time with a long, continuously proceeding
541 structuration process. On the contrary, SS-AAS reaches its initial setting with an abrupt
542 increase in the structuration process. At their initial setting times, the NH-AAS and SS-AAS
543 mixtures release similar heat and reach a similar reaction degree.
- 544 • At initial setting times, the C-A-S-H and N-A-S-H gels are both formed in the NH-AAS and
545 SS-AAS mixtures. The difference, however, is that the amount of C-A-S-H gel is relatively
546 similar to N-A-S-H gel in NH-AAS, while SS-AAS contains much more N-A-S-H gel.
- 547 • With time elapsing, more N-A-S-H gel in the SS-AAS mixture precipitates up to the end of
548 the induction period. However, the 3D N-A-S-H gel degrades to a chain-like C-A-S-H gel
549 after the acceleration peak recorded in the isothermal calorimetric curves.

550

551 The findings presented in this study can help researchers understand, to some extent, the early gel
552 composition and alkaline activation process of AAS mixtures. However, some remaining challenges
553 still need to be addressed in further research. For example, modeling the N-A-S-H structure could help
554 understand the reaction process. In addition, more research on the relationships between the
555 composition factors and inter-particle forces could also help better understand the interaction
556 mechanisms before setting times.

557

558 **Authorship contribution statement**

559 **Xiaodi DAI:** Methodology, Conceptualization, Investigation, Writing – original draft; **Serdar**
560 **AYDIN:** Methodology, Conceptualization, Writing – review editing; **Mert Yücel YARDIMCI:**
561 Methodology, Validation, Writing – review editing; **Gunter REEKMANS:** Investigation, Validation;
562 **Peter ADRIAENSENS:** Investigation, Validation; **Geert De SCHUTTER:** Funding acquisition,
563 Supervision, Writing - review & editing.

564

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571

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