Composition of C-S-H in pastes with increasing levels of silica fume addition

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10 Abstract

11 New results show that the microstructure development of cement-silica fume blends is very different 12 from plain cement. portlandite (CH) tends to precipitate as platelets and even around clinker grains as 13 "CH rims" and is consumed by pozzolanic reaction with silica fume. The Ca/Si ratio in the inner 14 product (IP) C-S-H decreases as CH is consumed to reach Ca/Si \approx 1.40-1.50 at the point when CH has 15 disappeared, and then drops down to 1.00 in absence of CH. At later ages, the IP C-S-H is often 16 composed of two distinct regions. The outermost (formed first) consists of originally high Ca/Si C-S-H, 17 which Ca/Si slowly decreases. The second (formed later) forms only once CH is no longer present and has a lower Ca/Si. Between 10 and 38°C, the main effect of increasing the temperature is to accelerate 18 19 the reaction of cement and increase the reactivity of silica fume. The changes in Ca and Si in the pore 20 solution of similar systems suggest that the composition of the solution and the solids reciprocally 21 influence each other.

- 22 Keywords: (B) Calcium silicate hydrate (C-S-H); (D) blended cement; (D) silica fume; (B) SEM; (B)
- 23 TEM

24 **1. Introduction**

The main hydrate phases in Portland cement pastes and blends of cement with supplementary
cementitious materials (SCMs) are calcium silicate hydrate (C-S-H) and portlandite (CH). The hyphens
in C-S-H indicate a variable composition. C-S-H in cementitious binders determines most of the
properties of the final material. They include the porosity, the chemical binding capabilities, and the

29 nature and amount of secondary phases, all of which affect the mechanical properties and durability. A

30 basic understanding of this phase, in particular in cement pastes, is essential to better understand the

31 impact of any change in the C-S-H on the final microstructure of cement pastes.

32 There are two main experimental approaches to study C-S-H. One approach is to synthesise it and 33 equilibrate it in dilute conditions (typically solution/solid ≈ 50), which allows the composition in 34 thermodynamic equilibrium to be established. The other is to characterise C-S-H within a cement paste 35 (with typical water/binder ratios of 0.3-0.7 [1]). Here it is formed by rapid dissolution-precipitation 36 processes but is not straightforward to analyse as it is part of a very heterogeneous microstructure. 37 C-S-H is often referred to as a "gel" as it has little or no long range order (peaks in XRD). However, 38 studies have shown that in both synthetic and paste systems it is made up of nanocrystalline regions 39 with an atomic structure resembling tobermorite and/or jennite [2]. In these minerals and in C-S-H, a 40 central layer of calcium and oxygen is flanked on both sides by tetrahedral silicate chains. These 41 "dreierketten" chains have a repeat unit of three. Two tetrahedra share an oxygen with the Ca-O layer 42 and the third is not directly bound to the Ca-O layer, but "bridges" adjacent paired tetrahedra. The 43 interlayer region contains variable amounts of water and other ions, notably calcium.

One very important characteristic of C-S-H is its variable composition, which is caused by defects and guest ions in its crystal structure. Several mechanisms have been described to allow variation of Ca/Si, although doubts still remain at Ca/Si over 1.50 [2]. It is known that the Ca/Si ratio of the C-S-H varies due to missing bridging silica tetrahedra [3], charge compensation of terminal oxygen atoms by calcium rather than protons and by the likely presence of calcium in the interlayer [4,5]. Aluminium may replace silicon in the bridging tetrahedra and, less commonly, in other sites [6,7]. SCMs generally supply more silica and alumina to the system, thus the final composition of the C-S-H and amount of secondary phases are affected this way. The (alumino-)silicate chains of C-S-H in blended cements condense to
form longer chains than non-blended systems [8,9].

53 The possible range of compositions is known for synthetic preparations. Solubility data in the 54 CaO-SiO₂-H₂O system reports a Ca/Si in C-S-H which can vary from ≈ 0.67 up to ≈ 2.00 depending on 55 the ionic activity of Si and Ca, generally with precipitation of portlandite occurring due to supersaturation of Ca with respect to the solid phase [1,4,10–12]. Pure C-S-H by most synthesis routes 56 57 only exists for Ca/Si = 1.50 or less, with most attempts to form C-S-H with higher Ca/Si causing the coprecipitation of portlandite [1,4,10-12]. Studies have reported changes in the structure of C-S-H 58 59 depending on the Ca/Si [3,12–15], on alumina substitution [16–19] and on the dependence of the composition of C-(A-)S-H (C-S-H with or without alumina substitution) to solution composition, 60 61 temperature and pH [20–23]. On the other hand, the C-S-H which forms with the hydration of plain 62 Portland cement is generally reported to have an average Ca/Si of approximately 1.75, e.g. [24] with 63 lower values generally reported for cement pastes containing supplementary cementitious materials 64 such as slag and fly ash [25–27]. We however lack a more precise idea about the range of possible 65 Ca/Si of C-S-H in pastes. We are therefore interested in exploring this using blends of Portland cement with low to high additions of silica fume. These systems are suitable for this purpose because of the 66 67 high reactivity of silica fume particles.

68 Silica fume is a highly reactive SCM which typically has a very small size – in the 100s of nm. Its 69 particles are mainly amorphous SiO₂. It is added in rather small amounts in practice, typically 5-10%, to 70 reduce bleeding, enhance mechanical properties and improve durability [28]. Higher dosages of silica 71 fume (up to 50%) are used however to investigate low-pH materials, for example [29].

Silica fume affects the hydration kinetics. It accelerates the hydration of C_3S because it provides more nucleation sites for hydrates to form and grow. This is one aspect of the "filler effect" which is the name given to the physical effect of blending cement with other materials without changing the chemical reactions of the system [30,31]. Silica fume has a particularly strong effect due to its small size. Silica fume can also react with CH to form additional C-S-H. Thermodynamic calculations for a cement containing increasing amounts up to \approx 50% by weight of silica fume [32], assuming complete reaction of both the clinker component and silica fume, show that increasing levels of replacement leads to a decrease in portlandite and the Ca/Si of C-S-H. The lower Ca/Si leads to an increase in alkali uptake which in turn can decrease the pH of the pore solution, which can bring additional changes to the microstructure (see Fig. 5 in [32]).

Transmission electron microscopy (TEM) has shown that the morphology of C-S-H in pastes is changed in presence of silica fume in blends hydrated at 40°C [33] to resemble foils which differ from the fibrils observed in PC.

85 Temperature does not appear to affect the Ca/Si in C-S-H from cement [34] nor its morphology [2,35].

86 An increase in temperature can change its density as seen by different grey levels and decrease the

bound water content [34]. In blended cements, temperature is expected to act on the kinetics of cement

88 and SCM reaction and can impact the Ca/Si and the Al/Si depending on the degree of reaction of the

89 SCM. The morphology of C-S-H does not appear to differ in blends either, e.g. [26].

In this paper new results on the C-S-H composition of real pastes are presented, measured over time in
pastes of water/binder = 0.4 by scanning electron microscopy with energy-dispersive spectrometry
(SEM-EDS) and compared to the amount of CH determined by thermogravimetric analysis (TGA). This
was done for temperatures within the range 10-38°C. The morphology of the outer product (OP) C-S-H
at later age was observed in several blends at 20°C by TEM.

95 **2. Materials and methods**

96 The blends studied are summarized in Table 1. They consisted of a plain Portland cement

97 (CEM I 32.5 R) with undensified silica fume (98.7% SiO₂). In two mixes quartz was also added to

98 increase the water to reactive binder (cement and silica fume) ratio without problems of bleeding. The

99 mixes with quartz were observed only at 90 days of hydration. The phase composition of cement is

100 given in Table 2. The chemical composition by XRF for all the materials is given in Table 3.

101 The cement and quartz were mixed in a Turbula T2F blender for 40 minutes. The silica fume was 102 dispersed in the distilled water using a hand blender for 2-4 minutes to form a slurry. It was set at 103 approximately 12,000 rpm. Each slurry was mixed for an additional 2 minutes just before each sample 104 was cast. The slurry was poured into the cement (or cement and quartz mixture) and mixed for two 105 minutes using a vertical mixer (IKA LABORTECHNIK RW20.n) with a rotation speed of about 1,600 106 rpm. This procedure was sufficient to limit the agglomeration of silica fume particles in the 107 microstructure. Also, all samples were workable without the need for superplasticiser. 108 Samples were cast into 50 ml plastic cylindrical tubes (35 mm diameter and 70 mm high) and then 109 stored first during 24 hours at 10, 20 or 38°C. They correspond to reasonable temperatures used in real 110 conditions and are not expected to destabilise Ettringite [36]. They were then removed from the mould 111 and stored in a second container of slightly larger width (38 mm diameter and 62 mm high). A small 112 amount of additional distilled water was added to maintain the sample under water. The amount of additional water was limited to minimize leaching of the pore solution. They were then stored again at 113 114 the same respective temperatures.

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Table 1: Mix design for the cement-silica fume blends

		Cement "PC" [wt%]	Silica fume "SF" [wt%]	Quartz "Q" [wt%]	SF/PC ratio [-]	/Water ra	(PC+SF) atio [-]	Water ra	:/binder atio [-]	
РС		100	-	-	0.00	0	0.40		0.40	
PC(Q) 9.1SF		81.8	9.1	9.1	0.11	0.44		0.40		
PC 10SF		90	10	- 0.11		0.40		0	0.40	
PC(Q)	20SF	60	20	20	0.33	0	.50	0	.40	
PC 258	SF	75	25	-	0.33	0	.40	0	.40	
PC 45SF		55	45	-	0.82	0.40		0	0.40	
	_							-		
	-	Table 2: Pha	se composition o	f the cement by	y XRD and I	Rietveld anal	ysis, in [wi	t%].		
	Alite	Table 2: Pha Belite	se composition o	f the cement by AF CaO	y XRD and D MgO	Rietveld anal	ysis, in [wissing SiO2	t%]. CaSO4	K ₂ SO	

	SiO ₂	Al ₂ O ₃	Fe ₂ O ₃	CaO	MgO	SO ₃	Na ₂ O ₃	K ₂ O	TiO ₂	P_2O_5	Mn ₂ O ₃	LOI
PC	21.1	5.6	2.6	64.9	1.6	2.8	0.2	0.9	0.3	0.1	0.1	1.9
Q	98.1	1.1	0.0	-	-	-	0.1	0.6	0.1	0.0	0.0	0.2
SF	98.7	0.3	0.0	0.2	0.0	-	0.1	0.3	0.4	0.0	0.0	0.5

At different hydration times slices approximately 2 mm thick were cut from the samples using a diamond saw and distilled water as lubricant. The slices were stored five days in isopropanol to remove the water by solvent exchange, with the solution being changed once within the first day to enhance the removal of water. After five days, the samples were stored in a desiccator for several days to evaporate the isopropanol.

Samples were ground by hand in a mortar for thermogravimetric analysis. TGA was carried out using a
Mettler-Toledo TGA/SDTA 851 balance with a 10°C/min ramp from 30°C to 900°C under a constant
30 ml/min flux of nitrogen.

129 For the preparation of SEM samples, a piece of dried paste was cut and impregnated in EPO-TEK 301 130 resin, prepolished using SiC paper (1200 grade) and isopropanol as lubricant, polished using several 131 diamond grades (9, 3 and 1 µm) with deodorised petrol as lubricant, and carbon coated with a layer 132 about 20-30 nm thick using a BAL-TEC CED 030 carbon evaporator. BSE images were taken using a 133 FEI Quanta 200 SEM equipped with a Bruker AXS XFlash 4030 EDS detector for point analysis. Only 134 measurements in IP rims larger than the estimated interaction volume of the electron beam (\approx 2-3 µm 135 diameter) were taken into consideration. The SEM operating voltage was 15kV. The spot size was 136 adjusted to yield a current of approximately 0.7 nA and an EDS livetime of about 3-4 seconds per point 137 analysis (50'000 total counts per spectrum). PB-ZAF with $\varphi(\rho z)$ matrix corrections were used to 138 quantify a preset list of elements (O, Na, Mg, Al, Si, P, S, Cl, K, Ca, Ti, Fe). The software used a 139 standard database comprised of several relevant oxides such as silica, alumina, jadeite and wollastonite 140 to determine the concentrations of each element. Oxygen was determined by stoichiometry.

In parallel work, it was shown [37,38] that the composition of C-S-H measured in SEM-EDS – when represented as a scatter plot (e.g. Si/Ca – Al/Ca) – could be estimated by choosing the value with the least intermixing of C-S-H with CH or other phases rich in calcium. Usually, the Ca/Si value is taken as the edge of the cloud of points (least intermixed points) in a plot of Al/Si against Si/Ca. In the samples studied here, most measurements of OP contained silica fume particles as well as C-S-H [38] within the interaction volume, particularly at high additions of silica fume. Therefore, only measurements from the
IP region were taken into consideration. Around 20 points were analysed for each sample. Values of
Ca/Si were taken as the average of these analyses and the error given is the standard deviation of these
points.

150 Thin lamellae for transmission electron microscopy (TEM) were prepared by taking smaller pieces of 151 dried paste and impregnating them using EMS Embed 812 resin. Each sample was then cut to a thin 152 slice (of dimensions 1.7 mm \times 1.7 mm \times 0.7 mm) using a WELL diamond wire saw, thinned down mechanically to a bevel by mean of the Tripod method (supplied by ALLIED Tech) to yield a thickness 153 154 of about 20-30 µm on the thick side. It was then glued to a copper ring, ion-thinned with an Argon PIPS (GATAN Precision Ion Polishing System Model 691) operated at maximum 1.5 keV to achieve electron 155 156 transparency and coated with a 5 nm layer of carbon using a CRESSINGTON 208 high vacuum carbon 157 coater. As a precaution, samples were coated only prior to observation in the TEM and were always 158 stored in a high vacuum desiccator. TEM observations were carried out using a FEI Tecnai Osiris at an 159 operating voltage of 80 kV in scanning TEM mode (STEM) with EDS mapping. A high spot size and 160 the nanoprobe mode were used to limit the beam current to less than 0.160 nA. Quantification was done 161 using the standardless Cliff-Lorimer ratio technique [39]. It ignores the effects of X-ray absorption and 162 fluorescence inside the material assuming the sample is thin enough. The background is subtracted from the peaks before concentrations C_A and C_B of elements A and B are calculated using the equation 163

164
$$\frac{C_A}{C_B} = k_{AB} \frac{I_A}{I_B}$$
(1)

165 This equation relates the above-background characteristic intensities I_A and I_B to the concentrations 166 using the factor k_{AB} which depends on the system and the accelerating voltage. Additional details on the 167 procedure can be found in [37] and in [38].

168 The heat of hydration was followed on a series of samples in a Thermometrics TAM Air isothermal

169 calorimeter set at 20°C and located in an air-conditioned room at approximately 20°C. For 10 g of

170 cement paste, 4.14 g of distilled water was used for each channel as a reference as this is of approximate

- 171 equivalent specific heat. The resulting heat release is normalised to the mass of the paste and the
- 172 anhydrous clinker content.

173 **3. Results and discussion**

174 **3.1. Kinetics**

- Figure 1 shows the calorimetry of the blends at 20°C. As widely reported, the reaction of C_3S is
- 176 strongly accelerated by the presence of silica fume [8,40-42]. The reaction of C₃A occurs earlier and is
- 177 enhanced [41] due to the adsorption of sulfate on C-S-H. Similar effects were observed at the other

temperatures, with an underlying acceleration in all reactions due to temperature.



179

180Figure 1: Isothermal calorimetry data for the four systems hydrated at 20°C. The heat release is divided by the181anhydrous cement content. 36 hours of hydration are shown.

182 Figure 2 shows the evolution of CH in the different blends. The rate of consumption increases with

183 temperature as expected.







Figure 2: %CH normalised to the clinker content for the 3 temperatures. (a) Data from the series hydrated at 10°C, (b) 20°C and (c) 38°C.

189 **3.2. Microstructure**

190 Figure 3 compares the microstructure of the blends at 20°C after 3 days of hydration. The addition of

- silica fume seems to accentuate the formation of gaps between the reacting grains and the hydrate shells
- 192 resulting in more and larger hollow "Hadley" grains from the hydration of small cement grains, as seen
- 193 for 25% and 45% substitution in Figure 3c and Figure 3d. This may be attributed to the fact that the
- 194 silica fume particles provide more sites for nucleation resulting in the formation of more "outer" and
- 195 less "inner" product at early ages. Similarly, the outer product regions appear more homogeneous in the
- 196 systems with silica fume, reducing the amount of large capillary pores which are still visible in the plain
- 197 cement (Figure 3a).



(b) PC-10SF





202

Figure 3: Pastes hydrated at 20°C for 3 days.

Silica fume also impacts the morphology of CH. In the plain cement, CH forms mostly as masses of CH with a few platelets (Figure 3a). In silica fume blends there are a majority of platelets. These changes in morphology are more pronounced as the level of silica fume increases. At the higher levels of silica fume replacement another interesting feature appears for some of the clinker grains. This is the formation of CH in the gap between the hydration shell and the reacting grain, an example of which is shown at higher magnification in Figure 4b. It appears that the presence of silica fume particles between

- the clinker grains makes the precipitation of calcium hydroxide in the outer space difficult (perhaps due
 to "poisoning" of CH growth [43]), such that the gaps become favourable sites for CH growth. Possible
 reasons for this are either that the silicate concentration in the gap is lower or that the available space
 favours the growth of portlandite.
 In the range of temperatures studied there was a relatively small impact of temperature on the
 appearance of the microstructures at similar degrees of hydration. In the plain cement system some
- 215 densification of C-S-H and coarsening of the porosity could be observed as previously reported [34]. In
- the silica fume blends such changes were less evident.







Figure 4: Microstructure evolution of PC 25SF blend hydrated at 20°C.

219 After 90 days of hydration many C-S-H rims around the clinker grains showed two distinct grey levels 220 an outer brighter rim (formed first) and a darker inner rim (formed later). They are referred to as "Early IP" and "Later IP". There was considerable variation in the rim thicknesses within any one sample, but 221 222 a general trend towards thicker darker parts with increasing level of silica fume replacement and 223 increasing temperature. Figure 5 illustrates this range of variation. Figure 6 shows a quantitative EDS 224 map of a two tone rim in the PC(Q) 20SF sample (10° C). Here it is clear that the calcium content (and 225 consequently Ca/Si ratio) in the darker inner part is lower, while the content of silica seems to be 226 similar. It should be noted that this kind of two tone rim is different from that formed in pastes hydrated a two different temperatures [44,45] where the change in grey level results from a change in density 227

rather than chemical compositions as here. The formation of the darker inner rims appears to coincide with the disappearance of calcium hydroxide as a separate phase in the microstructure, at which point a change in either one or both the Si and Ca concentrations in the pore solution will occur. This is supported by two recent examples of analyses of pore solutions from blends with 40% replacement of cement by silica fume where the Si in solution reached a maximum of 1.5 mmol·1⁻¹ [46] and 1.04 mmol·1⁻¹ [47]. This is around ten times greater than the typical concentration seen in a plain cement paste [48].



235

236Figure 5: Microstructure of matured samples (hydrated for 90 days) which illustrate the presence of two IP rims of C-237S-H.

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SEM-EDS map of PC(Q) 20SF (10°C)





Figure 6: (a) BSE SEM image of an area of the PC(Q) 25SF (10°C) sample analysed by SEM-EDS to map the composition of both the "Early IP" and the "Later IP". (b) The Ca/Si ratio quantified over the whole area. The Early IP has a slightly higher Ca/Si than the Later IP. The bright line between the Alite and Later IP is an edge effect due to the interaction volume. (c) and (d) show the quantified Ca and Si maps respectively.

3.3. C-S-H composition



249 1.25 ± 0.17 (28 days at 38°C) and in the 45% SF blend the lowest value of Ca/Si was 1.00 ± 0.05 (28 250 days at 38°C). Figure 7a shows the Ca/Si of the Early IP C-S-H plotted against the amount of CH 251 relative to the anhydrous cement content for all the samples. In Figure 7b the data for the blends 252 hydrated at 20°C are extracted to show how the Ca/Si values decrease over time. It can be seen that 253 there is a slow decrease to around \approx 1.40-1.50 while CH remains in the sample. When all the CH is 254 consumed there is a rapid decrease to Ca/Si = 1.00. 255 The lower limit for Ca/Si in presence of CH is broadly consistent with numerous experiments on synthesised C-S-H where the maximum Ca/Si before also precipitating CH is similarly around 256 257 1.40-1.50 [1,4,10–12]. It can also be seen that there is a tendency towards lower Ca/Si ratios in the

blends containing quartz which had a higher water to cementitious (clinker plus silica fume) ratio. This might indicate that greater availability of space facilitates the removal of calcium from the C-S-H as also seen in the work of Bazzoni [49] and Muller [50]. It could also be that the quartz improves the dispersion of silica fume particles due to increased shearing forces, therefore increasing the specific surface area in contact with the pore solution. Silica fume particles would thus be more reactive in

263 presence of quartz particles.



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Figure 7: Comparison between the Ca/Si in the Early IP C-S-H and the %CH normalised to clinker content. (a) The data from all series. (b) PC series hydrated at 20°C. The points are connected to show the decrease of Ca/Si with time.

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3.4. C-S-H morphology

269 TEM was used to study the morphology of the OP C-S-H in the pastes hydrated at 20°C for 90 days

270 (Figure 8). Unfortunately, no IP regions from the silica fume blends could be observed due to the small

- area thinned for analysis. In the plain cement the morphology of OP C-S-H with Ca/Si = 1.89 is
- 272 predominantly fibrillar (Figure 8a) as previously reported [2,51]. The foil like morphology is
- 273 predominant in the blends with silica fume (PC 10SF in Figure 8b and PC(Q) 25SF in Figure 8c). This
- is compared with the morphology of a C-S-H [52] synthesised at 20°C from CaO and SiO₂

(water/solid = 45) in a Nitrogen atmosphere and dried with ethanol according to a protocol detailed in [53]. It can be observed that the foil-like morphologies are very similar. It is seen here that the foil-like morphology can exist even in practical paste systems with only 10% silica fume. It is also seen that foillike morphologies exist in this series with Ca/Si = 1.41 or less. It does not however signify that the composition is the only factor determining the morphology.



(a) Plain cement (PC), 90 days, 20°C



(b) PC 10SF, 90 days, 20°C







Values of Ca/Si in the IP C-S-H (measured by SEM) and the OP C-S-H (measured by TEM) are compared in Figure 9 for the PC, PC 10SF and PC(Q) 20SF series. The Ca/Si in the plain cement was $\approx 1.80-1.90$ (taken in SEM from the edge of the cloud of points, not the average), in both the IP and OP. In the series with silica fume, the OP Ca/Si was systematically about 0.2 lower than the Ca/Si of the IP. This is not unexpected as the OP regions are closer to the silica fume particles which supply silica to the system.



290

291 Figure 9: Ca/Si of the C-S-H measured by EDS in the IP (by SEM) and the OP (by TEM). 292 It is interesting to note that despite the Later IP C-S-H being in close proximity to calcium-rich alite, the 293 C-S-H from the Later IP region has a lower Ca/Si then the Early IP. While the formation of C-S-H is 294 affected by the proximity of reactants, it is also affected by the pore solution. The pore solution in plain 295 cement generally remains calcium-rich, which is why in the plain cement, the IP and OP composition 296 are roughly the same. In the silica fume blends, the pore solution is calcium rich at first during the 297 formation of the Early IP but becomes poor in calcium and enriched in silica over time. Then the Later 298 IP C-S-H forms with lower Ca/Si. Over time, the pore solution therefore appears to dominate the formation of C-S-H. 299

300 **4. Conclusions**

301	The current study presents new results on the microstructure and C-S-H composition of PC-silica fume
302	blends hydrated at 10, 20 and 38°C and with low to high substitution levels.
303	• The growth morphology of CH is strongly affected by the presence of silica fume which
304	promotes the formation of platelets and even sometimes rims of CH around alite grains.
305	• It was observed that two rims of IP C-S-H exist in such blends. There was an outer rim ("Early
306	IP") forming first, followed by a darker grey rim ("Later IP") forming at a later stage when the
307	CH was depleted.
308	• The Ca/Si of the Early IP C-S-H decreases to \approx 1.40-1.50 as CH is consumed by silica fume. It
309	can drop down to 1.00 in absence of CH.
310	• The OP C-S-H morphology is foil-like in silica fume blends even with only 10% addition.
311	These observations suggest the following. At early times, when CH is still present, Ca-rich C-S-H
312	forms. At later age, when more SiO ₂ has reacted (and less CH is present), Si-rich C-S-H forms but the
313	Ca/Si ratio of the earlier formed C-S-H also decreases. The loss of calcium from the C-S-H is driven by
314	the consumption of calcium by the reaction of silica fume. It occurs relatively gradually while solid CH
315	is present to replenish calcium to the pore solution and then decrease dramatically when solid CH has
316	all been consumed. This corresponds to changes in the pore solution reported elsewhere [46,47] where
317	a decrease in the Ca concentration and pH is observed, and where an increase of the Si concentration is
318	also observed. This indicates that the composition of the solution and that of the solids reciprocally
319	influence each other.

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326 **6. References**

- 327 [1] Taylor, H.F.W., Cement Chemistry, 2nd ed., Thomas Telford, 1997.
- 328 [2] I.G. Richardson, Tobermorite/jennite-and tobermorite/calcium hydroxide-based models for the
- structure of C–S–H: applicability to hardened pastes of tricalcium silicate,[beta]-dicalcium silicate,
 Portland cement, and blends of Portland cement with blast-furnace slag, metakaolin, or silica fume,
 Cem. Concr. Res. 34 (2004) 1733–1777.
- [3] P. Yu, R.J. Kirkpatrick, B. Poe, P.F. McMillan, X. Cong, Structure of calcium silicate hydrate C–
 S–H Near-, Mid-, and Far-Infrared Spectroscopy, J. Am. Ceram. Soc. 82 (1999) 742–748.
- [4] A. Nonat, X. Lecoq, The structure, stoichiometry and properties of C–S–H prepared by C3S
 hydration under controlled conditions, in: Nucl. Magn. Reson. Spectrosc. Cem.-Based Mater.,
 Springer, Berlin, 1998: pp. 197–207.
- 337 [5] A. Nonat, The structure and stoichiometry of C–S–H, Cem. Concr. Res. 34 (2004) 1521–1528.
- [6] I.G. Richardson, G.W. Groves, The incorporation of minor and trace elements into calcium silicate
 hydrate (C–S–H) gel in hardened cement pastes, Cem. Concr. Res. 23 (1993) 131–138.
- [7] M.D. Andersen, H.J. Jakobsen, J. Skibsted, Incorporation of aluminum in the calcium silicate
 hydrate (C–S–H) of hydrated Portland cements: a High-Field 27Al and 29Si MAS NMR
 investigation, Inorg. Chem. 42 (2003) 2280–2287.
- [8] Z.-Q. Wu, J.F. Young, The hydration of tricalcium silicate in the presence of colloidal silica, J.
 Mater. Sci. 19 (1984) 3477–3486.
- [9] D.H. Justnes, Kinetics of reaction in cementitious pastes containing silica fume as studied by 29Si
 MAS NMR, in: D.P. Colombet, P.H. Zanni, D.A.-R. Grimmer, P.P. Sozzani (Eds.), Nucl. Magn.
 Reson. Spectrosc. Cem.-Based Mater., Springer Berlin Heidelberg, 1998: pp. 245–268.
- [10] S.A. Greenberg, T.N. Chang, Investigation of the colloidal hydrated calcium silicates. II. Solubility
 relationships in the calcium oxide-silica-water system at 25°, J Phys Chem. 69 (1965) 182–188.
- [11]X. Cong, R. Kirkpatrick, Si MAS NMR study of the structure of calcium silicate hydrate, Adv.
 Cem. Based Mater. 3 (1996) 144–156.
- [12] J.J. Chen, J.J. Thomas, H.F.W. Taylor, H.M. Jennings, Solubility and structure of calcium silicate
 hydrate, Cem. Concr. Res. 34 (2004) 1499–1519.
- [13] I. Klur, B. Pollet, J. Virlet, A. Nonat, C-S-H structure evolution with calcium content by
 multinuclear NMR, in: D.P. Colombet, P.H. Zanni, D.A.-R. Grimmer, P.P. Sozzani (Eds.), Nucl.
 Magn. Reson. Spectrosc. Cem.-Based Mater., Springer Berlin Heidelberg, 1998: pp. 119–141.
- [14]K. Garbev, M. Bornefeld, G. Beuchle, P. Stemmermann, Cell dimensions and composition of
 nanocrystalline calcium silicate hydrate solid solutions. Part 2: X-Ray and thermogravimetry study,
 J. Am. Ceram. Soc. 91 (2008) 3015–3023.
- 360 [15]K. Garbev, G. Beuchle, M. Bornefeld, L. Black, P. Stemmermann, Cell dimensions and
- 361 composition of nanocrystalline calcium silicate hydrate solid solutions. Part 1: Synchrotron-based
- 362 X-ray diffraction, J. Am. Ceram. Soc. 91 (2008) 3005–3014.
- [16] P. Faucon, A. Delagrave, C. Richet, J.M. Marchand, H. Zanni, Aluminum incorporation in calcium
 silicate hydrates (C–S–H) depending on their Ca/Si ratio, J. Phys. Chem. B. 103 (1999) 7796–7802.
- [17]G. Renaudin, J. Russias, F. Leroux, F. Frizon, C. Cau-dit-Coumes, Structural characterization of C S-H and C-A-S-H samples-Part I: Long-range order investigated by Rietveld analyses, J. Solid
 State Chem. 182 (2009) 3312–3319.
- [18]G. Renaudin, J. Russias, F. Leroux, C. Cau-dit-Coumes, F. Frizon, Structural characterization of C S-H and C-A-S-H samples-Part II: Local environment investigated by spectroscopic analyses, J.
 Solid State Chem. 182 (2009) 3320–3329.
- [19]X. Pardal, I. Pochard, A. Nonat, Experimental study of Si-Al substitution in calcium-silicate hydrate (C-S-H) prepared under equilibrium conditions, Cem. Concr. Res. 39 (2009) 637–643.
- [20] P. Faucon, T. Charpentier, A. Nonat, J.C. Petit, Triple-Quantum two-dimensional 27Al Magic
 Angle Nuclear Magnetic Resonance study of the aluminum incorporation in calcium silicate
 hydrates, J. Am. Chem. Soc. 120 (1998) 12075–12082.
- [21] R. Barbarulo, H. Peycelon, S. Leclercq, Chemical equilibria between C–S–H and ettringite, at 20 and 85 °C, Cem. Concr. Res. 37 (2007) 1176–1181.

- [22] I. Garcia Lodeiro, A. Fernández-Jimenez, A. Palomo, D.E Macphee, Effect on fresh C–S–H gels of
 the simultaneous addition of alkali and aluminium, Cem. Concr. Res. 40 (2010) 27–32.
- [23] X. Pardal, F. Brunet, T. Charpentier, I. Pochard, A. Nonat, Al-27 and Si-29 solid-state NMR
 characterization of calcium-aluminosilicate-hydrate, Inorg. Chem. 51 (2012) 1827–1836.
- [24] I.G. Richardson, The nature of C–S–H in hardened cements, Cem. Concr. Res. 29 (1999) 1131–
 1147.
- [25] I.G. Richardson, G.W. Groves, Microstructure and microanalysis of hardened cement pastes
 involving ground granulated blast-furnace slag, J. Mater. Sci. 27 (1992) 6204–6212.
- [26] A.V. Girão, I.G. Richardson, R. Taylor, R.M.D. Brydson, Composition, morphology and
 nanostructure of C–S–H in 70% white Portland cement-30% fly ash blends hydrated at 55 °C.,
 Cem. Concr. Res. 40 (2010) 1350–1359.
- [27] F. Deschner, B. Lothenbach, F. Winnefeld, J. Neubauer, Effect of temperature on the hydration of
 Portland cement blended with siliceous fly ash, Cem. Concr. Res. 52 (2013) 169–181.
- [28]T.C. Holland, United States, Federal Highway Administration, Silica fume user's manual, Federal
 Highway Administration, Washington, D.C., 2005.
- [29] B. Lothenbach, G. Le Saout, M. Ben Haha, R. Figi, E. Wieland, Hydration of a low-alkali CEM
 III/B–SiO2 cement (LAC), Cem. Concr. Res. 42 (2012) 410–423.
- [30] W.A. Gutteridge, J.A. Dalziel, Filler cement: The effect of the secondary component on the
 hydration of Portland cement: Part I. A fine non-hydraulic filler, Cem. Concr. Res. 20 (1990) 778–
 782.
- [31]E. Berodier, K. Scrivener, Understanding the filler effect on the nucleation and growth of C-S-H, J.
 Am. Ceram. Soc. (2014).
- [32]B. Lothenbach, K. Scrivener, R.D. Hooton, Supplementary cementitious materials, Cem. Concr.
 Res. 41 (2011) 1244–1256.
- [33]G.W. Groves, S.A. Rodger, The hydration of C3S and ordinary Portland cement with relatively
 large additions of micro silica, Adv. Cem. Res. 2 (1989) 135–140.
- 404 [34]E. Gallucci, X. Zhang, K.L. Scrivener, Effect of temperature on the microstructure of calcium
 405 silicate hydrate (C-S-H), Cem. Concr. Res. 53 (2013) 185–195.
- 406 [35] A.V. Girão, I.G. Richardson, C.B. Porteneuve, R.M.D. Brydson, Composition, morphology and
 407 nanostructure of C-S-H in white Portland cement pastes hydrated at 55 °C, Cem. Concr. Res. 37
 408 (2007) 1571–1582.
- [36] B. Lothenbach, T. Matschei, G. Möschner, F.P. Glasser, Thermodynamic modelling of the effect of
 temperature on the hydration and porosity of Portland cement, Cem. Concr. Res. 38 (2008) 1–18.
- [37] J.E. Rossen, K.L. Scrivener, Optimisation of SEM-EDS to determine C-S-H composition in matured cement paste samples, in preparation for Mater. Characterisation. (2015).
- [38] J.E. Rossen, Composition and morphology of C-A-S-H in pastes of alite and cement blended with
 supplementary cementitious materials, EPFL Thesis n°6294, 2014.
- [39] D.B. Williams, C.B. Carter, Transmission Electron Microscopy: A Textbook for Materials Science,
 Springer Science & Business Media, 2009.
- [40] W. Kurdowski, W. Nocuń-Wczelik, The tricalcium silicate hydration in the presence of active
 silica, Cem. Concr. Res. 13 (1983) 341–348.
- [41] H. Cheng-yi, R.F. Feldman, Hydration reactions in portland cement-silica fume blends, Cem.
 Concr. Res. 15 (1985) 585–592.
- [42] C.M. Dobson, D.G.C. Goberdhan, J.D.F. Ramsay, S.A. Rodger, 29Si MAS NMR study of the
 hydration of tricalcium silicate in the presence of finely divided silica, J. Mater. Sci. 23 (1988)
 4108–4114.
- [43]Z.-Q. Wu, J.F. Young, Formation of Calcium Hydroxide from Aqueous Suspensions of Tricalcium
 Silicate, J. Am. Ceram. Soc. 67 (1984) 48–51.
- [44] C. Famy, K.L. Scrivener, A.K. Crumbie, What causes differences of C-S-H gel grey levels in backscattered electron images?, Cem. Concr. Res. 32 (2002) 1465–1471.
- [45] K.L. Scrivener, Backscattered electron imaging of cementitious microstructures: understanding and
 quantification, Cem. Concr. Compos. 26 (2004) 935–945.
- [46] T.T.H. Bach, C.C.D. Coumes, I. Pochard, C. Mercier, B. Revel, A. Nonat, Influence of temperature
 on the hydration products of low pH cements, Cem. Concr. Res. 42 (2012) 805–817.

- 432 [47] B. Lothenbach, D. Rentsch, E. Wieland, Hydration of a silica fume blended low-alkali shotcrete cement, Phys. Chem. Earth Parts ABC. 70-71 (2014) 3-16. 433
- 434 [48] B. Lothenbach, F. Winnefeld, C. Alder, E. Wieland, P. Lunk, Effect of temperature on the pore solution, microstructure and hydration products of Portland cement pastes, Cem. Concr. Res. 37 435 (2007) 483-491. 436
- [49] A. Bazzoni, Study of early hydration mechanisms of cement by means of electron microscopy, 437 438 EPFL Thesis n°6296, 2014.
- [50] A. Muller, Porosity characterisation across different cementitious binders by a multi-technique 439 approach, EPFL Thesis n°6339, 2014. 440
- 441 [51] I.G. Richardson, G.W. Groves, Microstructure and microanalysis of hardened ordinary Portland 442 cement pastes, J. Mater. Sci. 28 (1993) 265-277.
- [52] E. L'Hôpital, B. Lothenbach, G. Le Saout, D. Kulik, K. Scrivener, Incorporation of aluminium in 443 calcium silicate hydrates, Cem. Concr. Res. (2014). 444 445
 - http://dx.doi.org/10.1016/j.cemconres.2015.04.007.
- 446 [53]É. L'Hôpital, Aluminium and alkali uptake in calcium silicate hydrates (C-S-H), EPFL Thesis n° 447 6389, 2014.
- 448