Multiscale understanding of tricalcium silicate hydration reactions

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Tricalcium silicate, the main constituent of Portland cement, hydrates to produce crystalline calcium hydroxide and calcium-silicate-hydrates (C-S-H) nanocrystalline gel. This hydration reaction is poorly understood at the nanoscale. The understanding of atomic arrangement in nanocrystalline phases is intrinsically complicated and this challenge is exacerbated by the presence of additional crystalline phase(s). Here, we use calorimetry and synchrotron X-ray powder diffraction to quantitatively follow tricalcium silicate hydration process: i) its dissolution, ii) portlandite crystallization and iii) C-S-H gel precipitation. Chiefly, synchrotron pair distribution function (PDF) allows to identify a defective clinotbermorite, $\text{Ca}_{11}\text{Si}_9\text{O}_{28}(\text{OH})_2.8.5\text{H}_2\text{O}$, as the nanocrystalline component of C-S-H. Furthermore, PDF analysis also indicates that C-S-H gel contains monolayer calcium hydroxide which is stretched as recently predicted by first principles calculations. These outcomes, plus additional laboratory characterization, yielded a multiscale picture for C-S-H nanocomposite gel which explains the observed densities and Ca/Si atomic ratios at the nano- and meso- scales.

Le Châtelier already established that Portland cement hydration starts by the dissolution of calcium silicate species in water from the most soluble silicate phase. This process is followed by the precipitation of complex poorly-crystalline calcium-silicate-hydrates (generically named C-S-H gel) and the crystallization of Ca(OH)$_2$, portlandite, see overall reaction (1)\textsuperscript{2}. C-S-H gel is the main hydrated component in Portland cement pastes, and it is the main responsible for the strength and durability of the resulting mortars and concretes. Alite, an impure form of tricalcium silicate Ca$_3$SiO$_5$, is the main phase present in Portland cements and it has a slightly variable composition due to element-substitutions\textsuperscript{2}. The hydration of any alite shows, in addition to an initial fast (minor) dissolution, three main stages with time: i) induction (also known as dormant period), ii) acceleration; and iii) deceleration\textsuperscript{3}. Similar kinetic profiles take place in various heterogeneous hydration processes, for instance mineral weathering\textsuperscript{4} and glass alteration\textsuperscript{5}. There are two main theories to explain this early-age hydration behaviour. The first is known as ‘protective layer’ and it consists in the precipitation of a C-S-H gel diffusion barrier on the surfaces of alite particles which density and adherence change with time. The second is known as ‘geochemical model’ and it is related to the alite dissolution mechanism evolving from etch pit formation to step retreat\textsuperscript{6}. Despite one century of focused investigations, the underlying mechanism(s) for such time evolution is still strongly debated\textsuperscript{7,8}.

$$\text{Ca}_3\text{SiO}_5 + 5.2\text{H}_2\text{O} \rightarrow 1.2\text{Ca(OH)}_2 + (\text{CaO})_4\text{SiO}_2(\text{H}_2\text{O})_4.0$$ (1)

The hydration reactions of alite, (i) dissolution of crystalline alite, (ii) precipitation of C-S-H gel, and (iii) crystallization of portlandite, have been thoroughly studied by many techniques including laboratory X-ray powder diffraction\textsuperscript{9-12}, calorimetry\textsuperscript{13-15}, small-angle neutron scattering\textsuperscript{16}, advanced electron microscopies\textsuperscript{17,18}, $^{29}$Si magic-angle-spinning nuclear-magnetic-resonance\textsuperscript{19,20}, and theoretical simulations\textsuperscript{21-23}. C-S-H gel has a nanocrystalline nature and so its understanding is very challenging\textsuperscript{24} which includes the relationship with the solution...
where it is equilibrated. There are many reviews addressing this component and we cite just the most relevant and recent ones. An updated mechanism of growth of C-S-H gel has been very recently proposed. It is also important to add that the hydration of alite is affected by the presence of other species. This has been very recently exemplified by the study of the influence of aluminates, added as NaAlO$_2$, on the hydration kinetics of alite which was studied by a multi technique approach including molecular dynamics simulations to investigate the dissolution step at the nanoscale. Many studies have shown that the C-S-H gel aggregates contain poorly-crystalline interconnected nanoparticles described as globules, disks and foils which enclose water within nanoreservoirs, known as gel pore water. This gel water can evolve with time with consequences in C-S-H ‘bulk’ density. It is worth noting that this water within the gel is different from the capillary pore water (also known as free water, FW) as it cannot be removed without altering the properties of the system. Concerning the atomic arrangement within the nanocrystalline component of the C-S-H gel aggregates, several experimental and theoretical techniques have concluded that defective clinoptilolite is the best available approximation. Crystalline tobermorite-14Å, Ca$_4$Si$_6$O$_{15}$(OH)$_2$.7H$_2$O, has a Ca/Si ratio and density of 0.83 and 2.19 g cm$^{-3}$, respectively. The corresponding values for crystalline tobermorite-11Å, Ca$_4$Si$_6$O$_{15}$(OH)$_2$.5H$_2$O, are 0.67 and 2.40 g cm$^{-3}$, respectively. These previous observations are striking as they are not in straightforward agreement with two well-established key bulk measured properties: i) the Ca/Si ratio on the C-S-H aggregates ranges between 1.6–2.0; and ii) the density of nanoglobules (gel pore water excluded) range 2.5–2.6 g cm$^{-3}$. Defective clinoptilolite could justify a Ca/Si atomic ratio close to 1.2–1.3, but not higher than that. Very fine intermixing of C-S-H gel with calcium hydroxide has been proposed from electron microscopy which it could explain an overall Ca/Si ratio ranging 1.6–2.0.

The aim of this work is to contribute to the understanding of both the formation and nanostructure of C-S-H gel. Firstly, we used synchrotron X-ray powder diffraction (and calorimetry) for in situ determining the dissolution of alite as well as the crystallization of portlandite and the precipitation of the gel. Secondly, we have employed synchrotron X-ray total scattering (and $^{29}$Si MAS-NMR) to study the short- and medium-range atomic arrangement in the C-S-H gel nanoparticles. With this knowledge and observations from electron microscopy and previous reports, we propose a model for this complex heterogeneous system, developing a multiscale picture (see Fig. 1) for the hydration of alite in order to explain the observed mass densities and Ca/Si atomic ratios at the different scales. At the nanoscale, below 10 nm, C-S-H gel are composed of a fine intermixing of defective clinoptilolite, particle sizes ranging 3–5 nm with Ca/Si ratio close 1.2, and monolayers of Ca(OH)$_2$. These aggregates generate the gel pores. At the mesoscale, between 10 and 100 nm, neat C-S-H gel appears relatively heterogeneous with (CaO)$_x$SiO$_y$(H$_2$O)$_z$ overall composition but slightly variable Ca/Si atomic ratios in different volumes. This is now explained by slightly different defective clinoptilolite to Ca(OH)$_2$-monolayers local ratios. At the microscale, above 100 nm, the hydration reaction of alite is well known resulting in crystalline Ca(OH)$_2$, also named portlandite, C-S-H gel and capillary water, see Fig. 1.

**Results**

**In situ calorimetric study.** The hydration reactions of tricalcium silicate were studied in situ by calorimetry and synchrotron X-ray powder diffraction (SXRPD). The particle size distribution (PSD) (diameter) for as-received alite was quite large, $D_{50} = 20.8 \mu$m, see Supplementary Fig. 1a. Therefore, two additional samples were prepared, see methods, with $D_{50} = 7.4$ and $2.7 \mu$m, Supplementary Fig. 1b,c, respectively. The values of $D_{10}$ and $D_{90}$ are also shown in Supplementary Fig. 1. The characterization of these three alites is given in the supplementary information. Heat flow calorimetry curves and cumulative heat released traces up to seven days are shown in Fig. 2a,b, respectively. Table 1 reports the key values obtained from the calorimetric study including the alite reaction degree that can be estimated as the overall heat of hydration for tricalcium silicate is known, 517 Jg$^{-1}$. This is an approximation as the impurities in alite can play a role as well as the structural defects. The four studied pastes are labelled as C$_{3}$S$_{21}$µm$_{045}$, C$_{3}$S$_{21}$µm$_{080}$, C$_{3}$S$_{7}$µm$_{080}$ and C$_{3}$S$_{3}$µm$_{080}$ to highlight their PSDs and water-to-alite mass ratios, i.e. 0.45 means a ratio of 0.45. C$_{3}$S$_{21}$µm$_{080}$-qz and C$_{3}$S$_{21}$µm$_{080}$-qz stands for samples with 10 wt% of quartz and their calorimetries were recorded for the sake of comparison with the SXRPD study where 10 wt% of quartz as internal standard was employed.

**In situ synchrotron X-ray powder diffraction study.** SXRPD data taken in capillaries of unaltered pastes were analyzed by Rietveld methodology employing the internal standard method, in this case quartz, for amorphous quantification. For this in situ study, the overall amorphous values encompass not only the C-S-H gel content but also the FW because the hydration reactions were not arrested. To a first approximation, the bound water (the crystallization water and the gel pore water) can be calculated according to reaction (1), as the amount of dissolved alite is known. Then, this calculated value is subtracted from the initial amount of water, in order to obtain the amount of FW for each hydration age. Finally, all the weight percentages were recalculated excluding the amount of FW in order to follow the evolution of the C-S-H gel component, which it includes the nanopore gel water and any amorphous calcium hydroxide.

Figure 3a shows the phase content evolution with time for C$_{3}$S$_{21}$µm$_{080}$ paste from SXRPD data. It is worth noting that, at 5 hours of hydration (which was our first measurement), alite was partly dissolved, $\approx$4 wt%, and $\approx$4 wt% of C-S-H gel had precipitated. The crystallization of portlandite started later, close to 7.5 hours. At 14 hours, only 2.8 wt% of crystalline portlandite was measured. The hydration reaction progressed slowly up to 100 days of hydration as it can be shown in Fig. 3a. As an example, Supplementary Fig. 2a shows the Rietveld plot for this paste at 14 hours of hydration. Finally, the quantitative phase analysis results obtained at 100 days (2400 h) was 11.8, 22.7 and 62.1 wt% of unreacted alite, crystalline portlandite and C-S-H gel, respectively. The remaining content, 3.4 wt%, was belite, Ca$_3$SiO$_4$. 
Figure 3b shows a similar study for C3S_7 µm_080. For this sample, the hydration kinetics is faster. At the first measurement time, 2 hours, 4.6 wt% of alite was already dissolved, with the precipitation of ≈ 4 wt% of C-S-H gel and the crystallization of ≈ 0.4 wt% of portlandite. At 14 hours, 10.4 wt% of crystalline portlandite was measured. Supplementary Fig. 2b shows the Rietveld plot for this paste at 14 hours of hydration. A transition from the accelerated hydration reaction kinetics to decelerated kinetics is clearly observed in the phase content evolutions close to 20 hours of hydration. At 100 days of hydration, 9.3 wt% of unreacted alite was measured coexisting with 25.3 wt% of portlandite and 62.8 wt% of C-S-H gel. The remaining content, 2.6 wt% was belite, Ca2SiO4.

The alite sample with the smallest PSD, see Supplementary Fig. 1c, was hydrated in order to minimize the amount of unreacted alite. The hydration was arrested, see methods section, in order to remove the FW, and this paste is labelled hereafter C3S_3 µm_080_arrested:16d. Rietveld quantitative phase analysis of LXRPD after 16 days of hydration, see Supplementary Fig. 3a, gave: 1.1 wt% of unreacted alite, 20.5 wt% of crystalline portlandite, 2.5 wt% of crystalline calcium carbonate and 75.9 wt% of amorphous content (mainly but not necessarily only C-S-H gel). Supplementary Fig. 3b shows the simulated X-ray diffraction pattern for the defective clinotobermorite T3_14sc structure, average particle size ≈ 5 nm, which has been used in the PDF study to fit the contribution of the nanocrystalline C-S-H gel. Supplementary Fig. 3c shows the simulated X-ray diffraction pattern for a single C-S-H nanoglobule composed by defective clinotobermorite and two monolayers of Ca(OH)2, at the nanoscale. (Bottom) Hydration reaction of tricalcium silicate at the nanoscale highlighting the three main components of colloidal C-S-H nanocomposite: nanocrystalline clinotobermorite, amorphous (monolayer) calcium hydroxide and gel pore water. The (approximate) densities, mass and volume percentages of the different components are also given for an overall water content of four water molecules per silicate.

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isolated Q₀ tetrahedra from unreacted alite. The silicate MCL (mean chain length) can be determined from the expression
\[ MCL = \frac{2(Q_1 + Q_2)}{Q_1} \]
The deconvolution of the spectrum shown in Fig. 4 gave 70.8% for Q₁ and 24.7% for Q₂. Therefore, MCL was 2.70. This value agreed well with previous reports for early age pastes. The ²⁹Si MAS-NMR spectra for two related samples gave very similar MCL values, see Supplementary Fig. 4. In addition, the determination of the average Ca/Si ratio is also an important parameter in any investigation of a C-S-H gel. Supplementary Fig. 5 shows a HRTEM micrograph with EDS data as an example. From 64 analysed points, the average Ca/Si atomic ratio was 1.75 ± 0.16. Supplementary Fig. 6 displays a FEGSEM micrograph (fracture cross-section).

**Total-scattering pair distribution function study.** The PDF data for C₃S₃µm_080_arrested:16d, see Fig. 5, have been analyzed using the same strategy previously reported by us. Initially, a high r-region, i.e.
Figure 3. Quantitative phase analysis results from SXRPD. (a) C₃S₂₁ μm₀₈₀ and (b) C₃S₇ μm₀₈₀. The lines for C-S-H gel (blue) and portlandite (green) show the theoretical amounts expected from the measured dissolution of alite according to reaction (1).

Figure 4. $^{29}\text{Si}$ MAS-NMR spectra for C₃S₃μm₀₈₀_arrested:16d measured. Spinning rate of 15 kHz and a magnetic field of 14.1 T.
40–70 Å, was analyzed with the contributions of the crystalline phases: portlandite and alite. The low amount of alite was computed as discussed in the supplementary information. The calcite content converged to zero. After refining all parameters, the final \( R_W \) value was 26.9%. The fit is displayed in Fig. 5a. The unit cell values for portlandite converged to \( a = 3.595 \text{ Å} \) and \( c = 4.918 \text{ Å} \). Secondly, all the previously determined parameters were kept fixed and the contribution from the nanocrystalline component of C-S-H gel was investigated in the \( r \)-region, from 10 to 25 Å. As it was previously reported, several crystal structures have been tested (Hillebrandite, Jennite, stoichiometric Tobermorite-14, stoichiometric Tobermorites-11 (monoclinic and orthorhombic), stoichiometric clinotobermorites, and selected structural descriptions from ref.42 which had a MCL close to 3.0) to fit the contribution of the nanocrystalline C-S-H gel, see Supplementary Table 1. Jennite structure led to the worst PDF fit as evidenced by its higher \( R_W \) value, see Supplementary Fig. 7. Hillebrandite has now been included as it has been very recently reported as a good model for the nanocrystalline component of C-S-H gel but it gave a poor fit to our PDF data. Supplementary Table 1 gives the \( R_W \) values for each PDF fit and the quantitative phase analysis results. The defective clinotobermorite structure, \( T3_14sc \) with \( \text{Ca}_{11}\text{Si}_9\text{O}_{28}(\text{OH})_2 \cdot 8.5\text{H}_2\text{O} \) composition, has been selected as it gave the best fit to the C-S-H gel nanoparticle contribution to the PDF profile. The unit cell values for the defective clinotobermorite \( T3_14sc \) structure converged to \( a = 11.255 \text{ Å}, b = 7.320 \text{ Å}, c = 42.415 \text{ Å} \) and \( \beta = 94.2^\circ \) and the isotropic ADPs were 0.0083 and 0.0160 Å\(^2\) for Ca, Si, respectively. The \( R_W \) was 27.7% and the final fit is displayed in Fig. 5b. For all the PDF fits, the ADP (U-thermal displacement parameter) value for O was not refined, the value was fixed to 0.070 Å\(^2\) which correspond to that of as received \( \text{Ca}_3\text{SiO}_5 \). The PDF fit gave the following quantitative phase analysis results: 2.2 wt% of anhydrous alite, 33.5 wt% of portlandite and 64.3 wt% of clinotobermorite.

Thirdly, the low \( r \)-region, 2 to 10 Å, was studied. The PDF fit based on the contributions of portlandite, alite and defective clinotobermorite resulted in a difference PDF curve, which was quite large, see Supplementary Fig. 8, especially in the very low \( r \)-region, suggesting an amorphous component. The scattering misfit closely corresponds to the theoretical PDF trace for an isolated monolayer of \( \text{Ca(OH)}_2 \) as recently suggested. Therefore, we have included a monolayer of \( \text{Ca(OH)}_2 \), atoms arranged as in portlandite, in the PDF fits, see supplementary information. The fit of the difference curve indicates that this monolayer \( \text{Ca(OH)}_2 \) is expanded along \( a \) and \( c \) directions, 3.8 and 5.3%, respectively and compressed along \( b \) direction, 2.1%. Overall, the volume of

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**Figure 5.** PDF refinements. Experimental (blue circles), fitted (red lines) and difference (grey lines) PDF patterns for C\(_3\)S\(_3\)\( \mu \)m\(_0\)80_arrested:16d (a) from 40 to 70 Å; (b) from 10 to 25 Å; and (c) from 2 to 10 Å. For details of the fits, the readers are referred to the text.
monolayer Ca(OH)$_2$ is expanded 7.0% when compared to the volume of crystalline portlandite. This volume expansion would lead to a density decrease from 2.24 g cm$^{-3}$ in bulk portlandite to 2.11 g cm$^{-3}$ in monolayer portlandite. The stretched monolayer portlandite structure was then used in the PDF refinement (Fig. 3c). As expected, the larger correlation in this refinement took place between the scale factors of monolayer-Ca(OH)$_2$ and crystalline-Ca(OH)$_2$. ~55%. The Rw value for the final fit, without computing alite, see Fig. 3c, dropped from 57 to 38% due to the contribution of monolayer portlandite.

**Discussion**

The hydration kinetic of as received alite was slow with the maximum in the heat flow curve close to 30 hours, with a reaction degree of ~9%, see Table 1 and Fig. 2. The presence of 10 wt% of crystalline quartz accelerates the hydration and the maximum takes place at 23 hours, with a reaction degree of 10.7% for the paste hydrated at a w/s = 0.80. This slow kinetics was due to the large average particle size, $\approx 20 \mu$m, and its relatively high iron content, 1.1 wt% expressed as Fe$_2$O$_3$. It is known that iron$^{31}$ and chromium$^{32}$ strongly delay alite hydration. Conversely, its kinetic is only slightly dependent on the water-to-alite ratio as previously reported$^{14,33}$. The calorimetric and SXRPD studies showed a strong acceleration of alite hydration for smaller particle size samples, see Table 1 and Figs 2 and 3, in agreement with previous reports$^{11,54}$. It is worth noting that the small peak evident in the heat flow curve at 5 h for C$_3$S$_7$µm$_{080}$, see Fig. 2, is very likely due to its bimodal PSD with a minimal fraction of very small particle size close to 0.6 µm, see Supplementary Fig. 1b. The acceleration due to the presence of quartz, known as ‘filler effect’$^{35}$, is much more pronounced for the alite sample with the largest particles. This observation point towards that alite dissolution is kinetically the limiting reaction.

We underline that the first 45 minutes of hydration is not recorded in our calorimetric study and this could neglect the measurement of the hydration of $\approx 5\%$ of alite, according to the results from the SXRPD study. In any case is worth comparing the reaction degrees from calorimetry (with quartz!), Table 1, and those obtained from the quantitative analysis of SXRPD data, Fig. 3. For C$_3$S$_{21}$µm$_{080}$ at 23 h (the time of the maximum of the heat flow curve), a reaction degree (as defined by the transformed fraction) of 11% (it could be $\approx 16\%$ taking into account the fast dissolution) is obtained from the calorimetric study and $\approx 19\%$ is obtained from the synchrotron study. For C$_3$S$_7$µm$_{080}$ at 20 h, a reaction degree of 21% (which it could be $\approx 26\%$) is obtained from the calorimetric study and $\approx 38\%$ is obtained from the synchrotron study. We justify this disagreement due to the main differences between the two set of experiments: i) the rotation of the capillary in the diffraction study with its associated shear effect;$^{15}$ and ii) the slightly higher temperature of the diffraction study when compared to the calorimetric study, see methods. It is key to compare results from different techniques but it is also important to understand the role of the different experimental conditions that sometimes are necessary to ensure the maximum attainable accuracy.

It has been reported from *in situ* laboratory powder diffraction data, Bragg-Brentano geometry, that the crystalline portlandite content was about one third smaller than that expected from alite dissolution according to reaction (1)$^{96,98}$. This is not the case in our synchrotron powder diffraction study as the used methodology (rotating the capillary in transmission and merging data from three capillary positions), improves the accuracy of the analyses. Figure 3 also displays the expected amounts of portlandite and C-S-H gel from alite dissolution which were in very good agreement with the measured values. Finally, it is worth noting that at very early hydration ages, the measured crystalline portlandite contents are slightly smaller than the expected ones. We speculate that this could be due to the initial precipitation of amorphous calcium hydroxide.

At 100 days, the RQPA results for C$_3$S$_{21}$µm$_{080}$ were 11.8, 22.7 and 62.1 wt% of unreacted alite, crystalline portlandite and C-S-H gel, respectively. The corresponding values for C$_3$S$_7$µm$_{080}$ were 9.3, 25.3 and 62.8 wt%, respectively. According to equation (1), the expected amounts of portlandite and C-S-H gel were 23.4 and 61.4 wt%, and 24.4 and 63.7 wt%, for C$_3$S$_{21}$µm$_{080}$ and C$_3$S$_7$µm$_{080}$, respectively. The good agreement between the determined and expected contents, for both samples, indicates the accuracy of the methodology and the suitability of the stoichiometric coefficients in reaction (1).

The PDF analysis of the total scattering data for C$_3$S$_{3}$µm$_{080}$ arrested:16d paste agrees well with our previous report$^{96}$. Moreover, the very small amount of unreacted alite, lower than 2 wt%, yielded a PDF study with less uncertainties. The crystal structure which gave the best fit to the PDF data in the region 10 to 25 Å was that of a staggered-chain clinotobermorite$^{42}$. More studies are needed to confirm if this structural description is the best option for the hydration of alite under different conditions and in Portland cements. Its average silicate chain length of 3.0 is in agreement with the MCL value, 2.7, obtained by $^{29}$Si MAS-NMR. However, its Ca/Si ratio, 1.22, does not agree with the Ca/Si ratio of the C-S-H determined by electron microscopy, 1.75, which is widely reported in bibliography$^{94,95}$ and previously confirmed by reaction (1). However, these Ca/Si ratios are not obtained at the same probing length scales. The Ca/Si ratio of defective clinotobermorite measured by PDF analysis is obtained at the nanoscale, probing scale: 1–3 nm. The Ca/Si ratio measured by HAADF-STEM is obtained at the mesoscale, probing scale $\approx 50$–100 nm. The Ca/Si ratios determined from PEQSEM and Rietveld refinement of SXRPD data are obtained at the microscale, probing scale $> 500$ nm. The apparent disagreement in the Ca/Si ratios can be reconciliated by the hypothesis that C-S-H gel aggregates arising from the hydration of alite, is a composite formed by a fine intermixing at the nanoscale of defective clinotobermorite nanoglobules, sizes $\approx 4$ nm, and amorphous Ca(OH)$_2$, size $< 2$ nm, see Fig. 1.

The overall multiscale picture for the hydration of alite is shown in Fig. 1 where reaction (1) at the microscale is depicted on top and its breakdown at the nanoscale is depicted at the bottom. The densities of the different components are also reported as well as their percentages (mass and volume). The hypothesis of existence of monolayer Ca(OH)$_2$ and its fine intermixing with defective clinotobermorite is supported in three ways. I. The PDF analysis in the 2–10 Å region improves notably with the inclusion of monolayer portlandite, $R_w$ decreases from 57
to 38%. Unfortunately, the analysis of the scales factors does not allow to obtain the weight percentage of the monolayer Ca(OH)_2 component. II. The determined expansion of monolayer Ca(OH)_2 in this study, 7.0%, is in agreement with the expansion recently predicted by first principles calculations when studying pure calcium hydroxide systems.\(^7\) This density functional theory theoretical work reported the stability of monolayer Ca(OH)_2 and an expansion of the average Ca-O bond from 2.36 to 2.38 Å from crystalline portlandite to monolayer Ca(OH)_2. III. The PDF analysis in the 10–25 Å length scale gave crystalline portlandite and nanocrystalline clinotbermorite contents of 33.5 and 64.3 wt%, respectively (see Supplementary Table 1). Our model reported at the bottom of Fig. 1, and rescaled to take into account 2 wt% of unreacted alite, gives 36.8 and 61.1 wt%, respectively. The relative close agreement between these two sets of values can be also interpreted as an indirect support of the model.

Conclusion
The in situ SXRPD study has confirmed the stoichiometry of alite hydration reaction to yield portlandite and C-S-H gel with \((\text{CaO})_{3.8}\text{SiO}_2(\text{H}_2\text{O})_{1.8}\) average composition. Chiefly, by using high-resolution synchrotron PDF analysis, it has been found that C-S-H gel is heterogeneous at the nanoscale being composed of defective clinotbermorite, with approximate composition \(\text{Ca}_{0.8}\text{Si}_0.2\text{O}_3(\text{OH})_2\cdot8.5\text{H}_2\text{O}\) and monolayers of Ca(OH)_2. With these results and observations from electron microscopy and previous reports, a multiscale model for the hydration of alite is proposed (see Fig. 1) which explains the observed mass densities and Ca/Si atomic ratios at the relevant scales. At the nanoscale, below 10 nm, C-S-H gel are composed of a fine intermixing of defective clinotbermorite, particle sizes ranging 3–5 nm with Ca/Si ratio close 1.2 and \(\rho\approx2.5\) g cm\(^{-3}\), and monolayers of Ca(OH)_2, \(\rho\approx2.1\) g cm\(^{-3}\). The calcium silicate component justifies the previously reported nanoglobules density, \(\rho\approx2.6\) g cm\(^{-3}\).\(^{3,39,37}\) These aggregates generate the gel pores. At the mesoscale, between 10 and 100 nm, neat C-S-H gel appears with variable compositions, Si/Ca ratio and water content, centred at \((\text{CaO})_{3.8}\text{SiO}_2(\text{H}_2\text{O})_{1.8}\). This is explained by slightly different defective clinotbermorite to Ca(OH)_2-monolayers local ratios and the variable gel pore water. Uneven water content also justifies the observed C-S-H gel densities at this scale, 1.9–2.1 g cm\(^{-3}\)\(^{32,6}\). At the microscale, above 100 nm, heterogeneous \((\text{CaO})_{3.8}\text{SiO}_2(\text{H}_2\text{O})_{1.8}\) gel and homogeneous portlandite are arranged enclosing volumes of water, termed capillary water. The picture reported here should be taken into account for developing theoretical models. Furthermore, synthetic C-S-H gels (with Ca/Si ratios < 1.4) may have very different properties as the monolayer calcium hydroxide component could be absent. Finally, the new model explains a striking feature of the hydration of cements blended with fly ash where portlandite content is measured to decrease much less than predicted by thermodynamic modelling. Although portlandite is still present, the Ca/Si ratio in C-S-H is observed to decrease from close to 1.8 in plain pastes to close to 1.4 in fly ash blends\(^{28}\). This is now explained by the consumption of Ca(OH)_2 monolayer component of the C-S-H gel in the pozzolanic reaction.

Methods
Full details about the Methods can be found in the supplementary information.

Sample preparation. Monoclinic tricalcium silicate, alite, was acquired from Mineral Research Processing M.R.PRO. Its chemical composition determined by XRF was: 72.3 wt% CaO, 25.5 wt% SiO_2, 1.1 wt% Fe_2O_3, 0.5 wt% MgO and 0.5 wt% Al_2O_3. For the in situ synchrotron X-ray powder diffraction studies, the anhydrous mixtures were mixed with 10.00 wt% of SiO_2 (99.5%, AlfaAesar) as an internal standard\(^{45}\).

\(\text{C}_3\text{S}_21\mu\text{m}_080\) labels the paste produced by using as received alite with a water-to-alite mass ratio of 0.80. \(\text{C}_3\text{S}_7\mu\text{m}_080\): The as received alite was milled in a vibratory mill (Retsch, mod. MM200) and the resulting powder was mixed with a water-to-alite mass ratio of 0.80. For the PDF study, as received alite was attrition milled and hydrated at a water-to-silicate mass ratio of 0.80 for 16 days at 20° C. Finally, the hydration stopped. The heat flow was collected up to 7 days at 20 °C. The sample was labelled \(\text{C}_3\text{S}_3\mu\text{m}_080\text{_arrested:16d}\). For the sake of comparison in the calorimetry studies, an Ordinary Portland Cement (OPC) from Financiera y Minera S.A. has been used. The samples have been labelled as OPC_045, OPC_080 and OPC_080_qz to indicate the w/s mass ratios. Furthermore, label _qz denotes the OPC with added quartz as standard.

Particle Size Distribution (PSD). Average particle size and particle size distribution for the alite samples were measured using a laser analyzer, Mastersizer S, Malvern, UK.

BET surface area. The specific surface areas of the selected samples were measured by multi-point N_2 adsorption with a BET (ASAP 2420, Micromeritics, USA) instrument.

Calorimetry. The isothermal calorimetric study was performed in an eight channel Thermal Activity Monitor (TAM) instrument using glass ampoules. The heat flow was collected up to 7 days at 20 °C.

Thermal analysis. Differential thermal analysis (DTA) and thermogravimetric (TGA) measurement for \(\text{C}_3\text{S}_3\mu\text{m}_080\text{_arrested:16d}\) was performed in a SDT-Q600 analyzer from TA instruments (New Castle, DE).

NMR study. \(^{29}\)Si MAS-NMR (Magic Angle Spinning Nuclear Magnetic Resonance) spectrum for \(\text{C}_3\text{S}_3\mu\text{m}_080\text{_arrested:16d}\) was recorded at RT on a Bruker AVIII HD 600 NMR spectrometer (field strength of 14.1 T) at 156.4 MHz. The Chemical shift was referenced to an external solution of tetramethylsilane.

Laboratory X-ray powder diffraction (LXRPD) with internal standard. LXRPD data for \(\text{C}_3\text{S}_3\mu\text{m}_080\text{_arrested:16d}\) was collected on a D8 ADVANCE (Bruker AXS) diffractometer (SCAI – Universidad de Malaga) equipped with a Johansson monochromator, using strictly monochromatic Mo-K\(_\alpha\) radiation, \(\lambda = 0.7093 \text{ Å}, \text{in transmission geometry (0/0)}\). Sample was mixed with 20 wt% of internal standard (\(\gamma\text{-Al}_2\text{O}_3\)).
Synchrotron X-ray powder diffraction (SXRPD). For the phase evolution study, SXRPD patterns were collected in Debye-Scherrer (transmission) mode using the X-ray powder diffraction endstation of BL04-MSPD beamline at ALBA synchrotron (Barcelona, Spain). The total acquisition time was 6 min per dataset. For the PDF study, SXRPD data for C$_3$S$_{3.5}$m$_{0.8} \_arrested:16d$ were collected for 3 h at the same diffractometer. The temperature inside the experimental hutch was 28°C.

Rietveld data analysis. Rietveld analyses were performed using the GSAS suite of programs and the EXPGUI graphic interface. The non-crystalline content (amorphous and nanocrystalline) was determined by the internal standard methodology.

Pair Distribution Function data analysis. PDF experimental data were obtained using PDFgetX$^{33}$ with $Q_{\text{max}} = 21 \text{ Å}^{-1}$. Quantitative phase analysis was obtained by using the PDFgui software$^{34}$ and CMI-diffpy complex modeling software$^{35}$.

Electron microscopy study. High resolution transmission electron microscopy (HRTEM) measurements were carried out using a FEI Talos F200X microscope equipped with X FEG and super-X EDS (Energy Dispersive Spectroscopy) system with four silicon drift detectors (SDDs) which operates at an accelerating voltage of 200 kV. FEGSEM (Field Emission Gun Scanning Electron Microscopy) micrographs and EDS analysis were performed in a Helios Nanolab 650 Microscope (FEI Company) with a retractable CBS Backscatter detector (annular solid-state device) and X-Max 50 mm$^2$ detector (Oxford instruments).

Data Availability. All synchrotron X-ray powder diffraction raw data files underlying this article can be accessed on Zenodo at https://doi.org/10.5281/zenodo.1027759, and used under the Creative Commons Attribution license.
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Acknowledgements
This work has been supported by Spanish MINECO through BIA2014-57658-C2-2-R, which is co-funded by FEDER, BIA2014-57658-C2-1-R and 13 (IEDI-2016-0079) grants. We also thank CELLS-ALBA (Barcelona, Spain) for providing synchrotron beam time at BL04-MSPD beamline.

Author Contributions
A.C. and M.A.G.A. designed the research; A.C., D.L.Z. and J.D.Z.G. carried out the sample preparations, I.S., I.S., J.D.Z.G., A.G.D.T., D.L.Z., S.G.S. and A.C. performed the laboratory characterization; O.V., A.C., A.G.D.T. and D.L.Z. carried out the synchrotron experiments; A.C. and M.A.G.A. analyzed the synchrotron data; M.D. contributed to the PDF analysis; A.C. and M.A.G.A. wrote the paper which was revised with contributions from all authors.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-26943-y.
Competing Interests: The authors declare no competing interests.
This supplementary information contains:

1. Extended Methods section.

2. Alite sample characterization.

3. Thermal analysis characterization for C₃S_3µm_080_arrested:16d.

4. Supplementary tables:

   **Supplementary Table 1.** Selected results for the synchrotron PDF analysis for C₃S_3µm_080_arrested:16d paste in the 10-25 Å r-region, using different structural descriptions for the nanocrystalline fraction of C-S-H phase.

5. Supplementary Figures:

   **Supplementary Figure 1.** Particle size distribution (diameter) and cumulative measured in volume, of the following materials (a) as received alite, (b) vibratory milled alite and (c) attrition milled alite.

   **Supplementary Figure 2.** SXRPD Rietveld plots at 14 hours of hydration for (a) C₃S_21µm_080 and (b) C₃S_7µm_080. The main peaks are labelled as follow: portlandite (●), alite (●) and added internal standard, SiO₂ (●).
Supplementary Figure 3. (a) LXRPD (Mo-Kα1 radiation) Rietveld plot for C₃S₃μm₀₈₀_arrested:16d. The main peaks are labelled as follow: portlandite (●), alite (●) and added internal standard, α-Al₂O₃ (●), (b) Simulated XRPD pattern for the defective clinotobermorite T3_14sc structure with particle size of approximately 5nm, using the same wavelength (Mo-Kα1 radiation) and (c) Raw SXRPD pattern for the PDF study of C₃S₃μm₀₈₀_arrested:16d paste. SXRPD pattern for the empty capillary is also shown (red line).

Supplementary Figure 4. ²⁹Si MAS-NMR spectra for (a) C₃S₃μm₀₈₀_arrested:16d, (b) C₃S₃μm₀₈₀_non-arrested:32d (a second preparation batch for the 3 μm alite sample) and (c) C₃S₁₃μm₀₈₀_arrested:34d. The Mean Chain Length values are depicted and the intensity of the Q₀ resonances indicates the unreacted alite fraction. Spinning rate of 15 kHz and a magnetic field of 14.1 T.

Supplementary Figure 4. High-angle annular dark-field scanning transmission electron (HAADF-STEM) micrograph for C₃S₃μm₀₈₀_arrested:16d. Three independent analyses obtained by EDS are also included as examples.

Supplementary Figure 5. Field emission gun scanning electron (FEGSEM) micrograph for C₃S₃μm₀₈₀_arrested:16d.

Supplementary Figure 6. Experimental (blue circles), fitted (red lines) and difference (grey lines) PDF patterns for C₃S₃μm₀₈₀_arrested:16d from 10 to 25 Å using (a) clinotobermorite T3_14sc and (b) Jennite. The arrows highlight interatomic distance features poorly fitted by the Jennite structural description.

Supplementary Figure 7. Experimental (blue circles) and fitted (red solid line) PDF patterns for C₃S₃μm₀₈₀_arrested:16d in the 2 to 15 Å r-range with three components: crystalline
Portlandite, unreacted alite and clinotbermorite. Difference curve is shown as a grey line which clearly shows the presence of an amorphous constituent. Simulated PDF curves for a monolayer (red) and a double layer (blue) calcium hydroxide, crystalline portlandite (green) and clinotbermorite T3_14sc (pink) are also included.

**Supplementary Figure 8.** Thermogravimetric data for C₃S_3μm_080_arrested:16d.

6. **Description of every synchrotron powder diffraction raw data set deposited open access.**

7. **References.**

1. **Extended Methods section.**

**Sample description.** Monoclinic tricalcium silicate, alite, was acquired from Mineral Research Processing M.R.PRO. Its chemical composition determined by XRF was: 72.3 wt% CaO, 25.5 wt% SiO₂, 1.1 wt% Fe₂O₃, 0.5 wt% MgO and 0.5 wt% Al₂O₃. For the *in situ* X-ray powder diffraction studies, the anhydrous mixtures were mixed with 10.00 wt% of SiO₂ (99.5%, AlfaAesar) as an internal standard. Pastes were prepared by mixing alite with water by hand in a small plastic baker for 1 min with a spatula and then immediately loaded into glass capillaries of 0.5 mm of diameter with a syringe. The capillaries were sealed with grease to avoid water loss. Capillaries which were prepared to be measured at ages of 7 days or later were kept inside closed plastic containers.

C₃S_21μm_080 labels the paste produced by using as received alite with a water-to-alite mass ratio of 0.80. C₃S_7μm_080: The as received alite was milled for 140 minutes, in cycles of 10 minutes with a rest-time between cycles of 10 minutes, in a vibratory mill (Retsch, mod. MM200). The resulting powder was mixed with a water-to-solid mass ratio of 0.80.

For the PDF study, as received alite was attrition milled (in house developed at UMA) with isopropanol for 4 hours in cycles of 10 minutes with a rest-time between cycles of 10 minutes and then dried at 100°C for 1.5 hours. Then, this alite was hydrated at a water-to-solid mass ratio of 0.80 for 16 days. The paste was poured into a hermetically closed Teflon® cylinder for 1 day. Subsequently, the cylindrical paste was taken out and stored within demineralised boiled water for
16 days at 20°C. Then, the paste was milled to fine powder in an agate mortar. Finally, to remove
the excess of water, the sample was filtrated in a Whatman system (90 mm diameter Whatman filter
with a pore size of 2.5 μm on a Teflon support) and washed twice with isopropanol and finally with ether\textsuperscript{2}. This powder was filled in a glass capillary of 0.7 mm of diameter. This sample was labelled
C\textsubscript{3}S\textsubscript{3} μm\textsubscript{080_arrested:16d}.

**Particle Size Distribution (PSD).** Average particle size and particle size distribution for the alite
samples were measured using a laser analyzer, Mastersizer S, Malvern, UK.

**Calorimetry.** The isothermal calorimetric study was performed in an eight channel Thermal
Activity Monitor (TAM) instrument using glass ampoules. Pastes were prepared *ex situ* by mixing
for 1 min ~3 g of each sample with the appropriated water, ~2 g, and they were immediately
introduced in the calorimeter. A stabilization period of 45 minutes was needed to start the
measurements. The heat flow was collected up to 7 days at 20°C.

**Thermal analysis.** Differential thermal analysis (DTA) and thermogravimetric (TGA)
measurement for C\textsubscript{3}S\textsubscript{3}μm\textsubscript{080_arrested:16d} was performed in a SDT-Q600 analyzer from TA
instruments (New Castle, DE). The temperature was varied from RT to 1000°C at a heating rate of
10 °C/min. Measurements were carried out in open platinum crucibles under nitrogen flow.

**NMR study.** \textsuperscript{29}Si MAS-NMR (Magic Angle Spinning Nuclear Magnetic Resonance) spectrum for
C\textsubscript{3}S\textsubscript{3}μm\textsubscript{080_arrested:16d} was recorded at RT on a Bruker AVIII HD 600 NMR spectrometer
(field strength of 14.1 T) at 156.4 MHz with a 2.5 mm triple-resonance DVT probe using zirconia
rotors at 15 kHz spinning rates. The experiment was performed with \textsuperscript{1}H decoupling (cw sequence)
by applying a single pulse (\(\pi/2\)), an excitation pulse of 5 μs, 30 s relaxation delay and 10800 scans.
The Chemical shift was referenced to an external solution of tetramethylsilane.

**Laboratory X-ray powder diffraction (LXRPD) with internal standard.** LXRPD data for
C\textsubscript{3}S\textsubscript{3}μm\textsubscript{080_arrested:16d} was collected on a D8 ADVANCE (Bruker AXS) diffractometer
(SCAI – Universidad de Malaga) equipped with a Johansson monochromator, using strictly
monochromatic Mo-K\textsubscript{α1} radiation, λ=0.7093 Å, in transmission geometry (0/0). Samples were
mixed with 20 wt% of $\alpha$-Al$_2$O$_3$ (AlfaAesar 42571) as internal standard. $\alpha$-Al$_2$O$_3$ was previously heated up to 1500°C for 20 hours and sieved <125 µm.

**Synchrotron X-ray powder diffraction (SXRPD).** For the phase evolution study, SXRPD patterns were collected in Debye-Scherrer (transmission) mode using the X-ray powder diffraction endstation of BL04-MSPD beamline at ALBA synchrotron (Barcelona, Spain)$^3$. The wavelength, 0.61878(3) Å, was selected with a double-crystal Si (111) monochromator and determined by using Si640d NIST standard (a=5.43123 Å). The diffractometer is equipped with a MYTHEN detector especially suited for time-resolved and extremely good signal-to-noise ratio experiments. The glass capillaries, 0.5 mm of diameter, were rotated during data collection at a speed of 100 rpm to improve diffracting particle statistics. To improve the accuracy of the results, three SXRPD patterns were collect at three different positions of every capillary and merged to produce the final dataset. The total acquisition time was 6 min per dataset (2 minutes per pattern) over the angular range 1-35° (20). The temperature inside the experimental hutch was 28°C.

For the PDF study, SXRPD data for C$_3$S$_3$µm$_080$ arrested:16d were collected for 3 h at the same diffractometer. The employed wavelength was 0.41236(1) Å and the glass capillary diameter was 0.7 mm. Five patterns were collected, each lasted 37 min, and merged in order to improve the signal-to-noise ratio in the large recorded angular range, 1 to 120° (20). No changes between individual patterns were observed.

**Rietveld data analysis.** Rietveld analyses were performed using the GSAS suite of programs and the EXPGUI graphic interface$^4$. Final global optimized parameters were: background coefficients, zero-shift error, cell parameters, and peak shape parameters using a pseudo-Voigt function. Portlandite and alite phases presented anisotropic lineshape broadening which was fitted by using the approach based of multidimensional distribution of lattice metrics$^5$. For the portlandite crystal phase preferred orientation was also optimized by employing the March–Dollase ellipsoidal preferred orientation correction algorithm$^6$. The non-crystalline content (amorphous and nanocrystalline) was determined by the internal standard methodology$^7$.$^8$. 

5
Pair Distribution Function data analysis. PDF experimental data was obtained using PDFgetX3\textsuperscript{9} with $Q_{\text{max}}=21$ Å\textsuperscript{-1}. Quantitative phase analysis information was obtained from the PDF data by using the PDFgui software\textsuperscript{10}. Final global optimized parameters were: scale factors, unit cell parameters and ADPs parameters. The delta2 value\textsuperscript{11,12} (low-r correlated motion peak sharpening factor) was fixed to 2 Å\textsuperscript{2}. The instrumental parameters were obtained by measuring a similar data set for crystalline nickel. Nickel PDF data analysis converged to $Q_{\text{damp}}=0.0030$ Å\textsuperscript{-1} and $Q_{\text{broad}}=0.0073$ Å\textsuperscript{-1}. The refinement of the difference curve (between 2 and 10 Å) was performed by means of CMI-diffpy complex modeling software\textsuperscript{13}, with a structural model constituted of a nanoparticle of single-portlandite, with no assumption of periodicity. Final optimized parameters were: scale factor, stretching in the three different crystallographic directions, and ADP parameter for Ca atoms. This refined nanoparticle was then used (and the structural parameters fixed), together with the crystalline phases, in the fit of the PDF (2-10 Å) again with CMI-diffpy.

Electron microscopy study. High resolution transmission electron microscopy (HRTEM) measurements were carried out using a FEI Talos F200X microscope equipped with X FEG and super-X EDS system with four silicon drift detectors (SDDs) which operates at an accelerating voltage of 200 kV. Spectra were collected in Scanning TEM (STEM) mode. The sample was placed on a 200 mesh copper grid coated with formvar and carbon. For the field emission gun scanning electron microscopy (FEGSEM) study, the selected samples (ground powder) were covered with iridium. FEGSEM micrographs and EDS analysis were performed in a Helios Nanolab 650 Microscope (FEI Company) with a retractable CBS Backscatter detector (annular solid-state device) and X-Max 50 mm\textsuperscript{2} detector (Oxford instruments). Backscattering electron imaging (BSEI) was performed at 5 kV acceleration, and EDS analysis at 10 kV. The software AZtec (v.1.0) was used to quantify.

2. Alite sample characterization.
SXRPD data of the anhydrous as received alite, C$_3$S$_{21 \mu m}$, was analyzed by Rietveld methodology employing the internal standard method for amorphous quantification. A mixture between the M$_1^{14}$ and M$_3^{15}$ tricalcium silicate polymorphs were needed to obtain the best fit.

The Rietveld quantitative phase analysis gave the following phase assemblage: 38.7(3) wt% of M$_1$-Ca$_3$SiO$_5$, 57.9(2) wt% of M$_3$-Ca$_3$SiO$_5$ and 3.4(2) wt% of $\beta$-Ca$_2$SiO$_4$. The amorphous content was negligible for this sample. The final unit cell parameters for the M$_1$-Ca$_3$SiO$_5$ converged to $a=9.2990(2)$ Å, $b=7.0847(1)$ Å, $c=12.1984(3)$ Å and $\beta=116.144(1)^\circ$ and for the M$_3$-Ca$_3$SiO$_5$ were $a=33.1289(8)$ Å, $b=7.0561(2)$ Å, $c=18.5844(4)$ Å and $\beta=94.247(2)^\circ$.

The same analysis was performed for the vibratory milled sample, C$_3$S$_{7 \mu m}$. The quantitative phase analysis results obtained were the following: 31.6(3) wt% of M$_1$-Ca$_3$SiO$_5$, 62.6(2) wt% of M$_3$-Ca$_3$SiO$_5$, 2.6(1) wt% of $\beta$-Ca$_2$SiO$_4$ and 3.2(1) wt% of amorphous content. The final unit cell parameters for the M$_1$-Ca$_3$SiO$_5$ were $a=9.2999(2)$ Å, $b=7.0857(2)$ Å, $c=12.1995(3)$ Å and $\beta=116.148(2)^\circ$ and for the M$_3$-Ca$_3$SiO$_5$ were $a=33.1302(9)$ Å, $b=7.0575(3)$ Å, $c=18.5840(5)$ Å and $\beta=94.32(2)^\circ$.

The Rietveld analysis for the attrition milled sample, C$_3$S$_{3 \mu m}$, was performed by using LXRPD data. The Rietveld quantitative phase analysis was the following: 26.8(8) wt% of M$_1$-Ca$_3$SiO$_5$, 44.6(9) wt% of M$_3$-Ca$_3$SiO$_5$, 2.4(2) wt% of $\beta$-Ca$_2$SiO$_4$ and 26.3(1) wt% of amorphous content. The final unit cell parameters for the M$_1$-Ca$_3$SiO$_5$ converged to $a=9.2900(8)$ Å, $b=7.0714(11)$ Å, $c=12.1878(17)$ Å and $\beta=116.157(8)^\circ$ and for the M$_3$-Ca$_3$SiO$_5$ were $a=33.051(17)$ Å, $b=7.066(2)$ Å, $c=18.526(4)$ Å and $\beta=94.42(4)^\circ$.

Finally, The PDF pattern of as received Ca$_3$SiO$_5$ was fitted with the M3 structure of Mumme $^{16}$ by using PDFGui. The obtained unit cell parameters and atomic displacement parameters (ADPs) were: $a=12.213$ Å, $b=7.085$ Å, $c=9.304$ Å and $\beta=116.1^\circ$; and 0.018, 0.011 and 0.070 Å$^2$ for Ca, Si and O, respectively. These values were used for the analysis of the minor fraction present in the sample C3S$_{3 \mu m}$$_{080}$$_{arrested:16d}$ paste and only the scale factor for alite was refined.
3. Thermal analysis characterization for C₃S₃μm_080_arrested:16d.

The thermogravimetric curve for C₃S₃μm_080_arrested:16d paste is shown in Supplementary Fig. 8. Several mass loss stages are observed in thermal trace. The mass loss from RT to 250°C, 17.7 wt%, is mainly ascribed to the water release from C-S-H gel aggregates. The weight loss measured between 250°C and 400°C, 2.4 wt%, could be partly related with the dehydration of amorphous calcium hydroxide. 5.7 wt% mass loss is measured between 400°C and 550°C, centered at 455°C, which corresponds to the water loss from crystalline portlandite. Two final weight losses are evident from 550 to 1000 ºC. The one observed close to 755°C, ≈2.0 wt%, is likely due to the CO₂ released of crystalline calcite. The other close to 660°C, ≈3.8 wt%, is very likely related to the CO₂ release from amorphous calcium carbonate and probably also to the release of organic solvent, used for the arresting of the hydration, absorbed in the C-S-H gel as previously reported17.

The components of the gel are nanocrystalline defective clinotobemorite, amorphous (monolayer) calcium hydroxide, and water within the nanopores. The weight losses for each component can be compared with that experimentally obtained for the C₃S₃μm_080_arrested:16d paste. For full reaction, the theoretical weight loss for the water gel pores, 13.27 wt%, plus the water in clinotobemorite, 5.28 wt%, sum 18.55 wt% which is comparable with the experimental weight loss between RT to 250°C, 17.7 wt% for this paste. Secondly, the calculated weight loss from Si-OH condensation and from amorphous Ca(OH)₂ is 0.62 and 3.24 wt%, respectively which can be compared with the water loss observed between 250°C and 400°C, 2.4 wt% plus 2.1 wt% from the contribution of the carbonation. Finally, the experimental water loss from crystalline portlandite, 5.7 wt% is compatible with the calculated value according to reaction (1), main text, 6.7 wt%. 
4. Supplementary tables:

**Supplementary Table 1.** Selected results for the synchrotron PDF analysis for C₃S₃₃μm_080_arrested:16d paste in the 10-25 Å r-region, using different structural descriptions for the nanocrystalline fraction of C-S-H gel.

| Phase                          | $R_w$ (%) | $Ca_3SiO_5$ (wt%) | Crystalline $Ca(OH)_2$ (wt%) | Nanocrystalline fraction of C-S-H (wt%) |
|-------------------------------|-----------|-------------------|-------------------------------|----------------------------------------|
| Hillebrandite (o), ICSD #80127| 33.4      | 3.7               | 55.6                          | 40.7                                   |
| Jennite (t), ICSD #151413     | 33.5      | 3.3               | 46.5                          | 50.2                                   |
| Tobermorite-14 (m), ICSD #152489| 33.1     | 2.9               | 44.5                          | 52.5                                   |
| Tobermorite-11 (o), ICSD #92941| 28.5     | 2.5               | 38.3                          | 59.1                                   |
| Tobermorite-11 (m), ICSD #87690| 28.7     | 2.4               | 36.9                          | 60.7                                   |
| Tobermorite-11 (o), ICSD #100405| 28.4    | 2.5               | 37.2                          | 60.4                                   |
| Clinotobermorite (m), ICSD #90036| 27.4    | 1.8               | 27.1                          | 71.1                                   |
| Clinotobermorite (t), ICSD #90034| 27.9    | 2.0               | 29.8                          | 68.3                                   |
| Clinotobermorite (m), T5_14sc* | 28.3     | 2.1               | 32.3                          | 65.5                                   |
| Clinotobermorite (m), T5_11sc* | 28.4     | 1.9               | 28.8                          | 69.3                                   |
| **Clinotobermorite (m), T3_14sc** | **27.7** | **2.2**           | **33.5**                      | **64.3**                               |

*Richardson, 2014; (m), (o) & (t) denotes monoclinic, orthorhombic and triclinic, respectively.
5. Supplementary Figures:

Supplementary Figure 1. Particle size distribution (diameter) and cumulative measured in volume, of the following materials (a) as received alite, (b) vibratory milled alite and (c) attrition milled alite.
Supplementary Figure 2. SXRPD Rietveld plots at 14 hours of hydration for (a) $C_3S_{21\mu m\_080}$ and (b) $C_3S_{7\mu m\_080}$. The main peaks are labelled as follow: portlandite (●), alite (▼) and added internal standard, SiO$_2$ (★).
**Supplementary Figure 3.** (a) LXRPD (Mo-Kα1 radiation) Rietveld plot for C₃S₃μm_080_arrested:16d. The main peaks are labelled as follow: portlandite (●), alite (▲) and added internal standard, α-Al₂O₃ (★), (b) Simulated XRD pattern for the defective clinotobermorite T3_14sc structure with particle size of approximately 5nm, using the same wavelength (Mo-Kα1 radiation) and (c) Raw SXRPD pattern for the PDF study of C₃S₃μm_080_arrested:16d paste. SXRPD pattern for the empty capillary is also shown (red line)
**Supplementary Figure 4.** $^{29}$Si MAS-NMR spectra for (a) C3S_3µm_080_arrested:16d, (b) C3S_3µm_080_non-arrested:32d (a second preparation batch for the 3 µm alite sample) and (c) C3S_13µm_080_arrested:34d. The Mean Chain Length values are depicted and the intensity of the Q$_0$ resonances indicates the unreacted alite fraction. Spinning rate of 15 kHz and a magnetic field of 14.1 T.
**Supplementary Figure 5.** High-angle annular dark-field scanning transmission electron (HAADF-STEM) micrograph for C₃S₃μm_080_arrested:16d. Three independent analyses obtained by EDS are also included as examples.

|        | Ca (%) | Si (%) | Ca/Si |
|--------|--------|--------|-------|
| Point 1 | 62.8   | 37.2   | 1.69  |
| Point 2 | 64.2   | 35.8   | 1.79  |
| Point 3 | 63.9   | 36.1   | 1.77  |

**Supplementary Figure 6.** Field emission gun scanning electron (FEGSEM) micrograph for C₃S₃μm_080_arrested:16d.
Supplementary Figure 7. Experimental (blue circles), fitted (red lines) and difference (grey lines) PDF patterns for C3S_3µm_080_arrested:16d from 10 to 25 Å using (a) clinotobermorite T3_14sc and (b) Jennite. The arrows highlight interatomic distance features poorly fitted by the Jennite structural description.
Supplementary Figure 8. Experimental (blue circles) and fitted (red solid line) PDF patterns for C₃S₃µm_080_arrested:16d in the 2 to 15 Å r-range with three components: crystalline portlandite, unreacted alite and clinotobermorite. Difference curve is shown as a grey line which clearly shows the presence of an amorphous constituent. Simulated PDF curves for a monolayer (red) and a double layer (blue) calcium hydroxide, crystalline portlandite (green) and clinotobermorite T3_14sc (pink) are also included.
Supplementary Figure 9. Thermogravimetric data for C₃S₃μₐₐ₉₈₉₀₈₉_arrested:16d.
6. Description of every synchrotron powder diffraction raw data set deposited open access.

All synchrotron X-ray powder diffraction raw data files underlying this article can be accessed on Zenodo at https://doi.org/10.5281/zenodo.1027759, and used under the Creative Commons Attribution license.

Files:

Synchrotron X-ray powder diffraction study for quantitative analysis

C3S_anh_ALL.dat: anhydrous C₃S_21µm
C3S_080_cap1_5h_ALL.dat: C₃S_21µm_080 at 5h of hydration
C3S_080_cap1_ALL_01_1707132228.dat: C₃S_21µm_080 at 5.5h of hydration
C3S_080_cap1_ALL_02_1707132300.dat: C₃S_21µm_080 at 6h of hydration
C3S_080_cap1_ALL_03_1707132331.dat: C₃S_21µm_080 at 6.5h of hydration
C3S_080_cap1_ALL_04_1707140003.dat: C₃S_21µm_080 at 7h of hydration
C3S_080_cap1_ALL_05_1707140034.dat: C₃S_21µm_080 at 7.5h of hydration
C3S_080_cap1_ALL_06_1707140105.dat: C₃S_21µm_080 at 8h of hydration
C3S_080_cap1_ALL_07_1707140137.dat: C₃S_21µm_080 at 8.75h of hydration
C3S_080_cap1_ALL_08_1707140208.dat: C₃S_21µm_080 at 9.25h of hydration
C3S_080_cap1_ALL_09_1707140239.dat: C₃S_21µm_080 at 9.75h of hydration
C3S_080_cap1_ALL_10_1707140311.dat: C₃S_21µm_080 at 10.25h of hydration
C3S_080_cap1_ALL_11_1707140342.dat: C₃S_21µm_080 at 10.75h of hydration
C3S_080_cap1_ALL_12_1707140414.dat: C₃S_21µm_080 at 11.25h of hydration
C3S_080_cap1_ALL_13_1707140445.dat: C₃S_21µm_080 at 11.75h of hydration
C3S_080_cap1_ALL_14_1707140516.dat: C₃S_21µm_080 at 12.25h of hydration
C3S_080_cap1_ALL_15_1707140548.dat: C₃S_21µm_080 at 13h of hydration
C3S_080_cap1_ALL_16_1707140619.dat: C₃S_21µm_080 at 13.5h of hydration
C3S_080_cap1_ALL_17_1707140650.dat: C₃S_21µm_080 at 14h of hydration
C3S_080_cap1_ALL_18_1707140722.dat: C3S_21µm_080 at 14.5h of hydration
C3S_080_cap1_ALL_19_1707140753.dat: C3S_21µm_080 at 15h of hydration
C3S_080_cap1_ALL_20_1707140825.dat: C3S_21µm_080 at 15.5h of hydration
C3S_080_cap1_17h_ALL.dat: C3S_21µm_080 at 17.5h of hydration
C3S_080_cap1_22h_ALL.dat: C3S_21µm_080 at 22h of hydration
C3S_080_cap1_24h_ALL.dat: C3S_21µm_080 at 24h of hydration
C3S_080_cap1_28h_ALL.dat: C3S_21µm_080 at 27.5h of hydration
C3S_080_cap2_36h_ALL.dat: C3S_21µm_080 at 36h of hydration
C3S_080_cap2_39h_ALL.dat: C3S_21µm_080 at 39h of hydration
C3S_080_cap1_42h_ALL.dat: C3S_21µm_080 at 42h of hydration
C3S_080_cap2_44h_ALL.dat: C3S_21µm_080 at 44h of hydration
C3S_080_cap1_48h_ALL.dat: C3S_21µm_080 at 48h of hydration
C3S_080_cap1_52h_ALL.dat: C3S_21µm_080 at 52h of hydration
C3S_080_7d_ALL.dat: C3S_21µm_080 at 7d of hydration
C3S_080_14d_rep_ALL.dat: C3S_21µm_080 at 14d of hydration
C3S_080_28d_ALL.dat: C3S_21µm_080 at 28d of hydration
C3S_080_3m_ALL.dat: C3S_21µm_080 at 3m of hydration
C3Sm_anh_ALL.dat: anhydrous C3S_7µm_080
C3Sm_080_cap3_2h_ALL.dat: C3S_7µm_080 at 2h of hydration
C3Sm_080_cap3_4h_ALL.dat: C3S_7µm_080 at 4h of hydration
C3Sm_080_cap1_ALL_01_1707132243.dat: C3S_7µm_080 at 5.75h of hydration
C3Sm_080_cap1_ALL_02_1707132315.dat: C3S_7µm_080 at 6.25h of hydration
C3Sm_080_cap1_ALL_03_1707132346.dat: C3S_7µm_080 at 6.75h of hydration
C3Sm_080_cap1_ALL_04_1707140018.dat: C3S_7µm_080 at 7.25h of hydration
C3Sm_080_cap1_ALL_05_1707140049.dat: C3S_7µm_080 at 7.75h of hydration
C3Sm_080_cap1_ALL_06_1707140120.dat: C3S_7µm_080 at 8.25h of hydration
C3Sm_080_cap1_ALL_07_1707140152.dat: C3S_7µm_080 at 8.45h of hydration
C3Sm_080_cap1_ALL_08_1707140223.dat: C3S_7µm_080 at 9.25h of hydration
C3Sm_080_cap1_ALL_09_1707140254.dat: C3S_7µm_080 at 10h of hydration
C3Sm_080_cap1_ALL_10_1707140326.dat: C3S_7µm_080 at 10.5h of hydration
C3Sm_080_cap1_ALL_11_1707140357.dat: C3S_7µm_080 at 11h of hydration
C3Sm_080_cap1_ALL_12_1707140429.dat: C3S_7µm_080 at 11.5h of hydration
C3Sm_080_cap1_ALL_13_1707140500.dat: C3S_7µm_080 at 12h of hydration
C3Sm_080_cap1_ALL_14_1707140531.dat: C3S_7µm_080 at 12.5h of hydration
C3Sm_080_cap1_ALL_15_1707140603.dat: C3S_7µm_080 at 13h of hydration
C3Sm_080_cap1_ALL_16_1707140634.dat: C3S_7µm_080 at 13.5h of hydration
C3Sm_080_cap1_ALL_17_1707140706.dat: C3S_7µm_080 at 14h of hydration
C3Sm_080_cap1_ALL_18_1707140737.dat: C3S_7µm_080 at 14.5h of hydration
C3Sm_080_cap1_ALL_19_1707140808.dat: C3S_7µm_080 at 15h of hydration
C3Sm_080_cap1_ALL_20_1707140840.dat: C3S_7µm_080 at 15.5h of hydration
C3Sm_080_cap1_18h_ALL.dat: C3S_7µm_080 at 18h of hydration
C3Sm_080_cap1_20h_ALL.dat: C3S_7µm_080 at 20h of hydration
C3Sm_080_cap1_24h_ALL.dat: C3S_7µm_080 at 24h of hydration
C3Sm_080_cap1_28h_ALL.dat: C3S_7µm_080 at 28h of hydration
C3Sm_080_cap2_35h_ALL.dat: C3S_7µm_080 at 35.5h of hydration
C3Sm_080_cap2_39h_ALL.dat: C3S_7µm_080 at 38.5h of hydration
C3Sm_080_cap1_42h_ALL.dat: C3S_7µm_080 at 42.5h of hydration
C3Sm_080_cap2_44h_ALL.dat: C3S_7µm_080 at 44h of hydration
C3Sm_080_cap1_48h_ALL.dat: C3S_7µm_080 at 48h of hydration
C3Sm_080_cap1_52h_ALL.dat: C3S_7µm_080 at 52h of hydration
C3Sm_080_7d_ALL.dat: C3S_7µm_080 at 7d of hydration
C3Sm_080_14d_ALL.dat: C3S_7µm_080 at 14d of hydration
C3Sm_080_28d_b_ALL.dat: C₃S₇μm_080 at 28d of hydration

C3S_milled_080_3m_ALL.dat: C₃S₇μm_080 at 3m of hydration

**Pair Distribution Function study for nanoscale analysis and characterization**

empty0p7_ALL.dat: empty capillary.

Ni0p7_ALL.dat: Nickel sample employed as standard.

C3S_080_milled_Marzo_ALL.dat: C₃S₃μm_080_arrested:16d.
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