Title
Synchrotron X-ray nanotomographic and spectromicroscopic study of the tricalcium aluminate hydration in the presence of gypsum

Permalink
https://escholarship.org/uc/item/6585m631

Journal
Cement and Concrete Research, 111

Author
Geng, Guoqing

Publication Date
2018-09-01

Peer reviewed
Synchrotron X-ray nanotomographic and spectromicroscopic study of the tricalcium aluminate hydration in the presence of gypsum

Guoqing Genga,b,*, Rupert J. Myersb,c, Young-Sang Yud, David A. Shapirod, Robert Winarskii, Pierre E. Levitzf, David A.L. Kilcoyned, Paulo J.M. Monteiroa,g

a Department of Civil and Environmental Engineering, University of California at Berkeley, Berkeley, CA 94706, United States
b Laboratory of Waste Management, Paul Scherrer Institut, Villigen PSI, Switzerland
c University of Edinburgh, School of Engineering, King’s Buildings, Sanderson Building, Edinburgh EH9 3FB, United Kingdom
d Advanced Light Source Lawrence Berkeley National Laboratory, Berkeley, CA 94720, United States
e Center for Nanoscale Materials, Argonne National Laboratory, Argonne, IL 60439, United States
f Advanced Light Source Lawrence Berkeley National Laboratory, Berkeley, CA 94720, United States
g Laboratory of Waste Management, Paul Scherrer Institute, 5232 Villigen PSI, Switzerland

ARTICLE INFO

Keywords:
- Tricalcium aluminate
- Hydration products
- Characterization
- X-ray diffraction
- Spectromicroscopy

ABSTRACT

The rheology of modern Portland cement (PC) concrete critically depends on the correct dosage of gypsum (calcium sulfate hydrate) to control the hydration of the most reactive phase - tricalcium aluminate (C3A). The underlying physio-chemical mechanism, however, remains unsolved mainly due to the lack of high-spatial-resolved and chemistry-sensitive characterization of the C3A dissolution frontier. Here, we fill this gap by integrating synchrotron-radiation based crystallographic, photon-energy-dependent spectroscopic and high-resolution morphological studies of the C3A hydration product layer. We propose that ettringite (6CaO·Al2O3·SO3·32H2O) is the only hydration product after the initial reaction period and before complete gypsum dissolution. We quantify the 2D and 3D morphology of the ettringite network, e.g. the packing density of ettringite at various surface locations and the surface dissolution heterogeneity. Our results show no trace of a rate-controlling diffusion barrier. We expect our work to have significant impact on modeling the kinetics and morphological evolution of PC hydration.

1. Introduction

Portland cement (PC) concrete is ubiquitous. The centrality of this technology to modern society is attributed to the low cost and global availability of the raw materials used in PC manufacture (mainly limestone and clay) and concrete production (water and aggregate, e.g., sand and stone), its robustness, controllability and high performance [1]. Precisely controlled rheology of fresh PC concrete is fundamentally important to produce high quality and durable PC concrete infrastructure, which is essential to reduce its environmental impact [1–4]. Gypsum (CaSO4·2H2O) addition is critical to control the fluidity and overall time window of flowability of fresh PC concrete mixtures to avoid ‘flash setting’ [1, 4–6] by retarding the hydration of the most reactive PC clinker phase – tricalcium aluminate (C3A).

The hydration of C3A can be delayed by minutes to hours depending on the C3A to calcium sulfate ratio in cement [1, 2, 5]. The resulting period of low reactivity enables a fresh concrete mixture to be workable; however, the chemistry underlying this mechanism has not been completely elucidated despite being researched for over half a century [5, 7–19]. This uncertainty has mostly persisted because direct evidence of this mechanism has not been reported. Two main hypotheses exist, which describe the inhibition of C3A hydration to occur via: (i) the formation of an ettringite (6CaO·Al2O3·SO3·32H2O) or/or a C-A-H (calcium aluminate hydrate) [8]; or (ii) the adsorption of Ca and/or S complexes (4CaO·Al2O3·SO3·18H2O, also denoted as s-AFm) [8] and/or hydroxyl AFm (OH-AFm, mainly 4CaO·Al2O3·19H2O (C4AH19)) [9, 10], which dehydrates to 4CaO·Al2O3·13H2O (C4AH13) at a relative humidity ≲ 88% [11, 12], and gradually converts to 3CaO·Al2O3·6H2O (C2AH6) over time [9]); or (ii) the adsorption of Ca and/or S complexes onto C3A surface sites [13–15]. The latter hypothesis has recently gained increasing support although the ‘diffusion barrier’ hypothesis has not yet been disproven, due mainly to ettringite precipitates hindering a direct observation of the C3A dissolution frontier [16, 17]. Therefore, a nanometer-resolved chemical-crystallographic probe is...
needed to more fully investigate the C₃A dissolution frontier. Conventional imaging methods, including scanning electron microscopy (SEM) [18] and transmission electron microscopy (TEM) [19], are restricted by destructive sample preparation and electron beam damage. Three-dimensional (3D) sub-micron scale characterization of the C₃A dissolution frontier has also never been reported [20].

Recent advances in synchrotron-based experimental technology have substantially improved the ability to obtain direct chemical and morphological information in hydrated C₃A systems: state-of-the-art soft X-ray ptychography imaging [21, 22] coupled with scanning transmission X-ray microscopy (STXM), and X-ray adsorption near-edge fine structure (XANES) [23] now enable transmission imaging at ~5 nm pixel resolution and chemical (energy) mapping at ~70 nm pixel resolution [24, 25], respectively. Here, we exploit this technology to reveal the mechanism by which C₃A hydration is retarded in the presence of gypsum (with a C₃A to gypsum mass ratio = 1:0.4, and a water-to-solid mass ratio = 10), aiming to identify features with sizes on the order of tens of nm at the C₃A dissolution frontier. These data are complemented by three-dimensional (3D) tomograms reconstructed from full-field transmission X-ray microscopy (TXM) data at a voxel resolution ~20 nm³ [26]. Crystallographic information is also obtained using in-situ wide angle X-ray scattering (WAXS). Small angle scattering (SAS) data is calculated from the X-ray ptychographic images [27, 28] and compared with in-situ small angle X-ray scattering (SAXS) results, to obtain quantitative morphological information of the reacting C₃A particle surface boundaries. The implications of these results on the contemporary understanding of the dissolution-retarding effect of gypsum on C₃A hydration, and on the computer-based cement hydration modeling are then discussed.

2. Experimental

2.1. Materials

Pure C₃A and hydrogarnet (C₃AH₆) powders were purchased from Mineral Research Processing (http://www.mineralresearchprocessing.fr/). Gypsum was purchased from Fisher Scientific (#S76764). The purities of these materials were verified by powder X-ray diffraction (XRD) (see SI for details). The C₃A powder has a size distribution from 1 μm to 100 μm, with d₅₀ ~ 20 μm [15]. Partially-hydrated C₃A samples were prepared by mixing gypsum, C₃A and ‘nапопуре’ water (1.82 x 10⁻² Ω cm, produced using a Barnstead NANOpure II with a filter size = 0.2 μm) in sealed plastic vials at room temperature (25 ± 2 °C). The samples were agitated frequently by hand and/or constantly by mechanical rotation. An initial mass ratio of C₃A to gypsum of 1:0.4 was used. A w/s = 10 was selected to fit the requirement of most experiments, and a w/s = 1 was also used for the X-ray ptychography experiments to verify the influence of sample dilution. Ettringite and s-AFm were independently prepared by precipitation from stoichiometric mixtures of C₃A, gypsum and water, following the reported methods [29]. The identities of these reference minerals were confirmed by XRD, as shown in the Supplementary Information (SI).

2.2. SEM

C₃A powder was spread on carbon tape pieces attached to steel SEM sample holders, which were then submerged into saturated gypsum-water solution. After certain hydration time (80s and 160s), the hydrated samples were then immediately rinsed with large quantities of 99.5% ethanol and dried in N₂ atmosphere to prevent carbonation. Samples were carbon coated before observation, using a Zeiss EVO® in second electron mode at 5 keV and 7 pA.

2.3. STXM and Al K-edge XANES

The STXM experiments were conducted at beamline 5.3.2.1 of the ALS (Advanced Light Source) of LBNL (Lawrence Berkeley National Laboratory) [21], which operates at 1.9 GeV and 500 mA using a bending magnet source and an active servo-stabilized toroidal pre-mirror that generates 10¹⁷ photons per second at the sample location. The resolving power of the radiation source is E/ΔE ~ 5000. Following the sample preparation method in previous work [30], partially hydrated C₃A samples prepared as described above were drop-cast onto 100 nm-thick Si₃N₄ windows (Norcada™), with excess solution removed using Kimwipes (Kimtech Science™). Samples were immediately transferred to the vacuumed STXM chamber. Single absorption contrast images were collected at an incident beam energy of 1575 eV (close to the Al K-edge). Stacks of absorption contrast images were recorded from 1552 to 1605 eV with step size 0.1 eV from 1565 to 1594 eV and step size 0.3 eV over the bordering central energy ranges of this investigation. A dwell time of 3 s was used. The image stack was aligned and analyzed with the aXis2000 software [31]. As reference minerals, hydrogarnet was dispersed with acetone whereas ettringite and s-AFm were dispersed with nanopure water, drop-cast and measured similarly. Internal calibration was conducted by calibrating the background absorption maximum to 1560 eV.

2.4. X-ray ptychography

X-ray ptychography imaging was conducted using the same STXM setup used for the Al K-edge XANES experiments. For these measurements, a 60 nm outer zone width zone plate focused a coherent soft X-ray beam (750 eV) onto the sample, which was scanned in 40 nm increments to ensure overlap of the probed areas. The effects of incoherent background signals were eliminated by implementing a background retrieval algorithm that iteratively determines a constant offset to the diffraction data. The sample was aligned with respect to the zone plate through an interferometric feedback system. A pixel size of 5 nm was used in image reconstruction. The efficient image resolution is determined to be ~17 nm by Fourier Ring Correlation (FRC; see SI for details) [32]. Small angle scattering calculations were performed using the ptychography images as input (the algorithm is available in SI).

2.5. In-situ WAXS and SAXS

In-situ WAXS and SAXS experiments were conducted at the WAXS/SAXS beamline of the Synchrotron Light Research Institute (SLRI), Thailand, which runs at 1.2 GeV and 75–125 mA. For the measurements, an incident beam of 9 keV was transmitted through two pieces of kapton™ film between which the sample was placed, encompassing samples from 2 to 120 min of hydration. The q-range covered was 0.04 to 42 nm⁻¹ for WAXS and 0.07 to 0.7 nm⁻¹ for SAXS, both with an angular step size of 0.25°.

2.6. TXM tomography

The TXM projections were collected at the Hard X-ray Nanoprobe Beamline, operated by the Center for Nanoscale Materials at Sector 26 of the Advanced Photon Source [26]. A cluster of partially-hydrated C₃A samples was mounted on a tungsten needle tip and placed on a multi-directional scanning stage. The incident beam was tuned to 9 keV and condensed by an elliptically shaped single-reflection glass capillary with a central stop. Transmission images were recorded while the sample was rotated from −90° to 90°. Ten images were collected at every 1° step, with an exposure of 10 s at each step to obtain better statistics. All tomographic 3D volume reconstructions were performed by an algebraic reconstruction technique (ART) with 20 iterations [33,34]; see details in SI.
reported to be poorly-crystalline AFm phases [9, 35]. The initial formed ettringite that formed later on. Synchrotron-radiation based in-situ WAXS, needles are short (Fig. 1b), but with similar diameter as the abundant et-

tingrite precipitates by 160 s (Fig. 1c). The initially formed particles within the here, from 2 to 120 min of hydration (Fig. 1d). The intensities of the major peaks of OH-AFm (e.g., C4AH19) at 10.7 Å, (8.3°, red arrow in Fig. 1d) [39], and s-AFm at 8.97 Å (9.9°, blue arrow in Fig. 1d) [40], are not observed during the first 120 min of hydration, indicating that the initially-formed AFm-type flakes, if initially formed, remain poorly crystalline, or are gradually decomposed with increasing hydration time. They do not seem to convert to C3AH6 (4.44, 3.35 and 3.14 Å (i.e. 20.1°, 26.6° and 28.4°, respectively, green arrows in Fig. 1d) [41]).

A previous synchrotron radiation XRD study of a similar system also revealed no trace of crystallized AFm-type phases before gypsum is completely consumed [42]. However, AFm-type crystals form immediately after mixing solid and water at a much lower initial gypsum-to-C3A ratio [10]. It seems that the crystallinity of the AFm-type phase depends largely on the dynamics of ion concentration change at the beginning of hydration.

Mass percentages of individual phases in the crystalline component of the reacted sample are calculated through Rietveld refinement (Fig. 1d); see details in SI. A quick initial dissolution of C3A is observed during the first 10 min. The dissolution of gypsum is slower with respect to C3A, since the percentage of gypsum clearly rises within the first 10 min. There is an initially observed ettringite formation (~3.5 wt %) at 2 min, which remains nearly unchanged within the first 10 min. Considering the low Al concentration in the liquid [16], the dissolved Al from C3A most likely precipitates as Al-bearing phases of low sulfur content, for instance AH3 and AFm. This is consistent with the observation of previous in-situ XRD study on cement hydration [43, 44].

After the first 10 min, a slow and steady rate of ettringite formation as well as gypsum and C3A dissolution is observed, until 120 min. Et-

tringite forms at a rate of ~2 wt% per hour with respect to the total crystalline solid mass, reaching 8 wt% at 120 min. After 120 min hy-

dration, the weight ratio of ettringite-to-C3A is 0.125 (Fig. 1e). The average diameter of C3A particles is ~20 μm [15]. If the surface of C3A particle is smooth, the volume of ettringite divided by C3A surface area is ~2.8 μm³/μm², using the existing density data [36, 37]. As shown in Fig. 1b, the formation of flake-like precipitates in the first minute precedes the significant precipitation of ettringite needles. Therefore, it is unlikely that they act as a physical barrier to retard the precipitation of ettringite within the first 2 min of hydration. However, it is still unclear whether an AFm-type layer persists as a physical barrier sur-

rounding C3A particles throughout the low-reactivity period, as it may be XRD-transparent, or masked by ettringite needles in the SEM images.

3.2. XANES Spectromicroscopic study

To further identify the solid phases, regardless of their crystallinity, on partly hydrated C3A surface, we collected the Al K-edge XANES spectra of eight regions parallel to the C3A dissolution frontier (BI-B8 in Fig. 2a,) using nanometer resolved STXM. For comparison, XANES spectra of pure solid phases relevant to the C3A-gypsum-H2O system were also measured and presented. In the Al K-edge XANES spectra,
peaks A, C and E correspond to the transitions of 1s to 3s-like, 3p-like and 3d-like states, respectively, and peak D corresponds to multi-scattering within adjacent neighbor shells [45]. The transition of the 1s to the 3s state is forbidden by the selection rule but is observable if the tetrahedral coordination is distorted and permits a mixture of Al s and p states, which is observed here. In good agreement with other Al-bearing minerals [45-48], the position of peak C, as shown in Fig. 2b, is a few eV lower for tetrahedral Al (Al[4]), i.e., in C3A [36], than for octahedral Al (Al[6]), i.e., in solid hydration products such as ettringite [37], s-AFm [40], C4AH12 [11] (dehydration of C4AH19 is likely to occur in vacuum conditions [12]) and C4AH11 [41]. The unique features of the reference spectra allow us to determine the identity of the solid hydration products.

In the Al K-edge XANES spectra of B1-B8, the intensities of the Al[4] peaks (blue short dashed lines) A (1568.6 eV), C (1572.5 eV), D (1579.3 eV) and E (1587.4 eV) shrink and the Al[6] peaks (red long dashed lines) A (1574.9 eV) and E (1585.9 eV) grow as functions of increasing distance away from the C3A particle (Fig. 2b). The spectra collected in regions B1 and B2 are almost identical to the Al K-edge XANES spectrum of C3A, which show strong intensities at the energies of its major tetrahedral Al (Al[4]) peaks A, C, D and E, indicating that these regions contain predominantly remnant C3A. The positions of the growing Al[6] peaks C and E in B1-B8 match with the corresponding peaks and positions for ettringite (1574.9 and 1585.9 eV respectively) rather than C4AH12 (1576.0 and 1588.0 eV respectively) or C4AH13 (1575.8 and 1588.2 eV respectively). Compared to the intensities and shapes of peaks C and A in the spectrum for ettringite, peak C (1574.7 eV) is significantly sharper and peak A (1569.4 eV) is much weaker in the spectrum of s-AFm. Therefore, regions B3-B6 contain significant amounts of C4A (Al[4]) and ettringite (Al[6]) and regions B7–B8 contain predominantly ettringite (Al[6]). After 150 min of hydration, ettringite is the only precipitate observed in the dissolution frontier of the C3A particle analyzed here.

The Al K-edge XANES of the full imaged region is well-fitted using the spectra of C3A (blue) and ettringite (green), with only a small residual signal (red) in the background region that is thus assigned to measurement noise (Fig. 2c). This result further demonstrates that ettringite is the only solid hydration product here at 150 min of hydration. There is no trace of AFm phases, although flake-like AFm dominate the surface morphology at initial hydration (Fig. 1b). There is also no trace of C4AH12 that is expected to precipitate from the destabilization of OH-AFm phases. This ettringite layer is ~1 μm thick on the bigger C3A particle (Fig. 2a, right), and slightly thinner on the smaller one (Fig. 2a, left).

3.3. Quantifying nano-morphology using X-ray ptychographic imaging and in-situ SAXS

X-ray ptychographic imaging is used to observe the morphology of the C3A dissolution frontier (Fig. 3). The image resolution is determined to be ~17 nm by Fourier ring correlation; see SI. Needle-like ettringite precipitates studied here are observed with similar thickness from 30 to 620 min of hydration in the water to solid mass ratio (w/s) = 10 system (Fig. 3a–c). The population of ettringite needles increases with increasing hydration time, but individual needles generally all grow to a maximum length of ~1 μm and do not seem to thicken significantly. Ettringite prefers to nucleate on C3A particles, and generally grow perpendicular to the C3A surface (dashed blue squares). The needle network is increasingly densified through interlacing of ettringite needles in confined regions (dashed yellow squares). By 620 min of hydration, the layer of ettringite needles are observed to directly contact the remnant C3A surface, i.e., no continuous physical barrier is observed in the interfacial zone (inset in Fig. 3c).

Fig. 3. Morphology of the C3A dissolution frontier in the w/s = 10 (Fig. 3a–c) and w/s = 1 (Fig. 3d–f) systems are similar. Ettringite needles reach a length of ~1 μm after 910 min of hydration and thicken negligibly in the w/s = 1 system. A solid ‘diffusion barrier’ is not identified at 910 min of hydration (Fig. 3f inset). C3A hydration kinetics is reported to be independent of the w/s between 1 and 10 [16, 49]. In both w/s systems, there exist continuous ‘gel-like’ areas (red arrows in Fig. 3a, d and e) on the particle surfaces that have distinct grey scale values compared to the remnant C3A and with dimension of a few hundred nanometers. They are observed to be abundant at a relatively early age of hydration, and intermixed with ettringite needles on the C3A surface. These gel-like morphologies are highly consistent with the flake-like AFm gel as shown in SEM images (Fig. 1b). However, this gel is not observed at a later age of hydration under the transmission microscope (Fig. 3b, c and f), indicating that the initially formed AFm flakes are unstable, and may dissolve to feed the precipitation of ettringite during the low-reactivity period. Therefore, the initially formed AFm gel is unable to act as the physical barrier described in the ‘diffusion barrier’ hypothesis to retard C3A dissolution throughout the low-reactivity period as stated in the introduction. This interpretation is consistent with the STXM and Al K-edge XANES results.

To quantify the morphology of the ettringite needles that precipitate under different degrees of spatial confinement, theoretical calculations of SAS (algorithm available in SI) are applied to the selected dashed blue (O1–O6, less confined) and yellow (C1–C2, more confined) square regions in Fig. 3, and are compared with in-situ SAXS measurements of
the w/s = 1 system (Fig. 4). The calculated SAS results generally resemble the in-situ SAXS results, and both can be interpreted using the form factor of a typical Guinier-Porod pattern of agglomerated isometric rods or cylinders [50]. This provides strong evidence that the needle-like morphology of ettringite at the C3A dissolution frontier is not an artefact of drying, which is suggested to alter the surface morphology of hydrated PC particles [51]. In a log-log plot of the intensity vs. scattering vector (q), the q1 and q_cutoff values define the lower and higher end, respectively, of a linear Porod region with slope between −4 and −3, corresponding to the surface fractal dimension. For q < q1, a nonlinear Guinier region with slope ranging from −1 to −3 is inferred, which corresponds to the volume fractal dimension. Due to the limitation of the SAXS experimental setup and the overall size of the selected ptychographic image regions (~1 μm² here), the low-q end of the Guinier region here is not small enough to reliably probe the length scale equal to or larger than the needle length (i.e., ≥1 μm).

As shown in Figs. 1c and 3, the needle-like ettringite precipitates are similarly sized. In an isometric rod model, the radius (r) of the rod can be related to q1 following Eq. (1) [50]:

\[ q_1 = \left(2d - 2\right)^{-d/r} \]

where d is the fractal dimension of the Porod region (further details regarding the interpretation of the SAXS data are presented in the SI), i.e., the absolute value of the slope when the logarithm of the intensity (base 10) is plotted as a function of q (Fig. 4a). The fitted needle radius, r, from the in-situ SAXS measurement increases slightly as a function of hydration time, i.e., from ~20 nm at 20 min of hydration to ~23 nm at 120 min of hydration, with a standard error of ~0.7 nm (Fig. 4b). A comparable result was obtained from the SAS data calculated from the ptychographic images at both w/s = 1 and 10, where the radii of ettringite crystallites was determined to be ~23 ± 5 nm (Fig. 4b). Needles in confined space (C1-C2) are of similar thickness to those growing in open space (O1-O6). Therefore, before complete dissolution of gypsum (i.e., ≤10 h), the typical radius of ettringite precipitates is < 30 nm, and is independent of the w/s and hydration time; similarly the aspect ratio of ettringite needles is generally around 40. As hydration progresses, the growth of ettringite crystals is mainly through new precipitates on the C3A surface, rather than the thickening or lengthening of individual ettringite needles.

3.4. Quantifying 3D morphology using nano-CT

To date, only 2D nano- and meso-scale morphologies of hydrated
C₃A have been reported, providing limited morphological information on the C₃A dissolution frontier. To further probe the 3D nano-scale morphology of the C₃A dissolution frontier, a full-field TXM tomographic study was conducted to yield a 3D image at a voxel resolution of 20 nm, as shown in Fig. 5, where the blue and green colors represent C₃A and ettringite, respectively. A movie of the volume rendering is presented in the SI. The segmentation of the background, ettringite and remnant C₃A was based on the reconstructed optical density (OD_recon) histogram of the 3D image. Further details of the procedures are in the SI.

As calculated from the segmented 3D image, the volume of the ettringite and remnant C₃A are ~40 μm³ and ~52 μm³, respectively. By simplifying the C₃A particles in Fig. 5 as 4 agglomerated particles with roughly spherical shape and smooth surface, the total surface area of C₃A is ~27 μm². The volume of ettringite to the calculated surface area of remnant C₃A is thus ~1.5 μm³/μm². This value is in the same order of magnitude as estimated using quantitative analysis of WAXS data, i.e. ~2.8 μm³/μm², and therefore suggests that the segmentation is reasonable. However, by directly counting the surface area using the segmented 3D image of C₃A, its surface area is ~249 μm², which is one magnitude larger than the estimated value by assuming a smooth C₃A surface. This discrepancy indicates that the partially dissolved C₃A surface is very rough.

The result shows that C₃A particles are non-uniformly reacted at 143 min of hydration (Fig. 5), with the solid hydration products on C₃A surfaces varying in thickness up to 1–1.5 μm, with some surface areas barely covered. This information is not observed in the 2D X-ray ptychographic images (Fig. 3) as they are transmission images. Heterogeneous C₃A dissolution is observed to generate sub-micron scale features, including highly reacted and rough surface regions (yellow arrows in Fig. 5c and d) and much smoother surfaces with low reaction extents (red arrows in Fig. 5c and d). There seems to be no intermediate status, i.e., the C₃A surface either dissolves extensively to form needle-like precipitations of constant maximum length (~1.5 μm), or stays almost unreacted. From a visual estimation, the highly dissolved C₃A surface area is about 40% of its total surface area at a hydration age of 143 min. In some low-reaction surface regions, the green colour layer is of single voxel thickness, which is not necessarily a thin layer of ettringite. In fact, the transmission images with higher resolution shown no evidence of such continuous layers (Fig. 3c and f). Therefore they are most-likely due to artifacts of the segmentation due to the voxels on the C₃A surface containing both solid and void regions.

Surfaces of inner cavities are also reactive. Any originally closed cavities participate in hydration as soon as the dissolution frontier exposes them to the solution. Non-uniform hydration is also observed in the inner cavities of remnant C₃A particles (Fig. 5c and d), some of which are more connected to the external particle surface than others. The amount of reactive C₃A surface is directly related to the speed of ettringite formation [16]. Here, the TXM results show that the significant proportion of C₃A surface reacts at very slow rate which highlights the non-uniform nature of C₃A surface dissolution and hydration, observed on both external surface and internal cavity surface.

Porosities of selected sub-volumes in Fig. 5a, V1 (solid hydration products grown in confined space), V2 (solid hydration products grown in open space) and V3 (remnant C₃A), are determined according to the attenuation law [52]. In the SI, we show that the porosity, φ, can be estimated using Eq. (2):

$$\phi = 1 - \frac{OD_{recon,mean}}{OD_{recon,max}}$$

(2)

where OD_{recon,mean} is the average reconstructed optical density of the studied sub-volumes, and OD_{recon,max} is the reconstructed optical density of a zero-porosity voxel of the studied phase (ettringite or C₃A); see further details in the SI. By applying this equation to the three sub-volumes, we determine the respective porosities of V1, V2 and V3 to be 53–67%, 73–81% and 20–38%. Therefore, after 143 min of hydration, the network of ettringite needles in open C₃A spaces (V2) is highly
porous, and the network of ettringite needles in confined spaces (V1) is denser but also significantly porous. The region representing remnant C3A (V3) contains non-zero porosity that may be caused by the presence of initial cavities.

The porosity of the cementitious binder critically influences its physical properties, and is therefore a key target of hydration modeling [1, 2]. In a randomly packed rod network, the volume fraction of solid rods decreases with their increasing aspect ratio, reaching ~10% when the aspect ratio is ~40 [53]. Here, the volume fractions of ettringite needles on C3A surface estimated using the 3D image is ~2–5 times the value in the randomly packed case, indicating that the ettringite needles are systematically oriented. Previous studies of ettringite-water suspension indicate that the ettringite volume fraction ϕ and the rod aspect ratio are related to various rheological properties such as suspension viscosity and the onset rate of shear thinning [54]. Therefore, this characterization of porosity and hydration product geometry on the sub-30 nm length scale, the first of its kind, can be used to develop and validate detailed 3D computational models of C3A hydration, nanostructure and rheology, a substantial step-change to a higher level of detail compared to existing formulations.

4. Discussion

Our results provide strong evidence that OH-AFm and s-AFm do not greatly retard the initial formation of ettringite as ettringite forms rapidly after a few minutes of hydration when most of the C3A surface is already covered by amorphous AFm phases. These AFm phases also do not form a C3A dissolution-inhibiting ‘diffusion-barrier’ in the low-reactivity period, because ettringite is identified by coupled STXM and Al K-edge XANES as the only precipitate on C3A particles after 150 min of hydration, but before complete gypsum dissolution. The initially formed AFm phases destabilize completely to ettringite during the low-reactivity period. At an imaging resolution of ~15 nm, no obvious ‘diffusion-barrier’ is observed at the C3A dissolution frontier after the initial hydration period. Our X-ray ptychography and TXM results also show that the ettringite needle network at the C3A dissolution frontier is highly porous. The porosities of this network are 53–67% and 73–81% in confined and open spaces after 143 min of hydration, respectively, meaning that it does not act as a physical barrier to retard C3A hydration, in contradiction to the ‘diffusion barrier’ hypothesis. Therefore, the results point toward an alternative mechanism of C3A dissolution retardation, e.g., by adsorption of Ca and/or S complexes onto C3A surface sites [8, 15, 55].

The morphology of the C3A dissolution frontier in the presence of gypsum in aqueous solution is similar in systems with w/s = 1 and 10. In both cases, ettringite precipitation predominantly occurs via the production of more individual surface-bound needles rather than the existing needles increasing in thickness and length. Ettringite needles with radii of ~20 nm are observed after 20 min of hydration and grow to ~25 nm thick by 15 h hydration; and the length of individual ettringite needles remain ~1 μm. The independence of the needle size on the w/s suggests that it is not the free space which determines the size of needle. The growth of ettringite needles may be determined by the solution chemistry facts close to nucleation sites, instead of the bulk liquid region far away from C3A surface. A maximum growth length is also observed for the hydration products of tricalcium silicate (C3S), calcium silicate hydrates, which is suggested to be merely determined by the solution chemistry [56].

On the other hand, the 3D image clearly indicates that the C3A surface either undergoes substantial dissolution to form needles of 1–1.5 μm length, or otherwise stays essentially unreacted. This heterogeneity indicates that C3A dissolution is also somewhat controlled by its crystalline surface chemistry and morphology [55]. Similar observation was reported recently for the hydration of a polished C3A surface [57]. The dissolution of mineral surfaces are often promoted by the surface concentration of defects, such as roughness, grain boundaries and dislocations [58]. As indicated by Fig. 5, the dominating surface defect seems to be heterogeneously distributed at the micron-scale, and varies from facets to facets of the C3A particles. This could be either the difference of roughness (amount of kinks) created during the grinding process, or the different cleavages (with different dissolution rate) created during the sintering process. However, we are unable to identify individual surface defects due to the limit of resolution.

Upon the above discussion, we now have a general picture of the studied reaction. When C3A surface is in contact with sulfur-containing water solution, it undergoes quick dissolution and formation of an AFm-type gel, followed by the low low-reactivity period. In the low-reactivity period, the AFm-type gel quickly vanishes, and C3A surface dissolution is very slow probably due to ion-complexation. Along with the slow dissolution, as soon as a defect site is exposed to solution, there is a localized quick dissolution of C3A, followed by the quick growth of ettringite until a certain size. In the surface regions where less defects exit, ettringite formation is hardly observed. In future work, two processes need urgent investigation in order to quantitatively predict the C3A hydration: 1) the dissolution of a single defect site on C3A surface, 2) the mechanism that ettringite only grows to a certain size.

For the first time, this paper provides a sub-30 nm 2D and 3D morphological and spectroscopic study of the C3A dissolution frontier in the presence of gypsum during the low-reactivity period. It advances the understanding of cubic C3A hydration in the presence of gypsum, through the central quantification of the chemistry and morphology of hydration product on C3A surface. This work also serves as a preliminary step to studying PC hydration with sub-30 nm resolution synchrotron radiation technologies. We expect that these techniques will provide key experimental evidence to validate multi-scale computational simulations of PC hydration down to the nano-scale [59–61], and also the macro-scale properties of fresh PC concrete [54, 62].

Acknowledgments

This work is funded by the Singapore-Berkeley Building Efficiency and Sustainability in the Tropics (SinBerBEST) Program. The authors acknowledge the support received from Ubonwan Khongpiboon, Panod Viseschitra and Thanakrit Chantra (Siam Cement, Thailand), and from Siriwat Soontaranon (SAXS beamline of Synchrotron light research institute, Thailand). The Advanced Light Source was supported by the Director, Office of Science, Office of Basic Energy Sciences, of the U.S. Department of Energy under Contract No. DE-AC02-05CH11321. Use of the Center for Nanoscale Materials and the Advanced Photon Source, both Office of Science user facilities, was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, under Contract No. DE-AC02-06CH11357. The authors want to acknowledge Jeffrey W. Bullard (NIST, USA) for the insightful discussion on C3A hydration. Guoqing Geng’s PhD study is supported by the Chinese Scholarship Council (file No. 201206090127).

Appendix A. Supplementary data

The supporting information available as a separate file, which includes the characterization of the raw materials, process of analyzing the WAXS and SAXS data, reconstruction and segmentation of the 3D TXM result, estimating the resolution of the ptychography images, and determining the peak positions of the XANES results. A video of the 3D rendering of the TXM result is also provided. Supplementary data to this article can be found online at https://doi.org/10.1016/j.cemconres.2018.06.002.

References
