The Hydration of β- and α’\textsubscript{H}-Dicalcium Silicates: An X-ray Spectromicroscopic Study

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KEYWORDS. Portland cement; dicalcium silicate; low carbon cement; silicate chain polymerization; calcium silicate hydrate; calcium coordination; morphology

ABSTRACT

Dicalcium silicate (C\textsubscript{2}S) is an important clinker mineral in Portland and belite-calcium sulfoaluminate cement. However, there is still a lack of information on the local degree of silicate polymerization and calcium coordination in C\textsubscript{2}S hydration products. This study aimed to
fill this gap by characterizing the hydration of two C$_2$S polymorphs, the $\beta$- and $\alpha'$$_H$ types, using scanning transmission X-ray microscopy coupled with ptychographic imaging. The results showed that the coordination of the Ca species in $\beta$- and $\alpha'$$_H$-C$_2$S had a distorted, cubic-like symmetry, whereas the Ca coordination of calcium silicate hydrate (C-S-H), the main hydration product, was structurally similar to that of tobermorite. The outer hydration product (Op) of both polymorphs exhibited an increasing degree of silicate polymerization over time. The inner hydration product (Ip) of $\beta$-C$_2$S polymerized slower than Op; however, the degree of silicate polymerization of both Ip and Op in the $\alpha'$$_H$-C$_2$S hydration system was comparable. The polymerization degree of Op was inhibited by adjacent $\beta$-C$_2$S particles, whereas in $\alpha'$$_H$-C$_2$S, the polymerization degree was more homogenous. Ptychographic imaging shows that the Op of $\alpha'$$_H$-C$_2$S exhibits coarser fibrils than the Op of $\beta$-C$_2$S, and a clear Op-Ip interface of hydrated $\beta$-C$_2$S is observed.

INTRODUCTION

Concrete is the second most consumed material after water $^1$. Its strength comes from the hydration of cement. The production of conventional Portland cement (PC) accounts for 8-9% of global CO$_2$ emission and 2-3% of energy consumption $^2$. The major compounds of PC clinker are tricalcium silicate (Ca$_3$SiO$_5$, also termed C$_3$S in cement chemistry notation), dicalcium silicate (Ca$_2$SiO$_4$, also termed C$_2$S; or belite as impure C$_2$S), tricalcium aluminate (Ca$_3$Al$_2$O$_6$), and tetracalcium aluminoferrite (Ca$_2$AlFeO$_5$). Among them, C$_3$S is the most abundant phase, but it has the highest environmental impact due to the high calcination temperatures and heavy CO$_2$ emission from decomposing limestone. The cement industry is under increasing pressure to reduce CO$_2$ emissions and energy consumption. Production of C$_2$S-rich PC at the expense of the
typically more abundant C\textsubscript{3}S is one way to lower the environmental impact of PC manufacturing. For example, belite-calcium sulfoaluminate (BCSA) cement is a very promising alternative to PC because the calcination temperature of BCSA is 200°C lower than PC\textsuperscript{4}. BCSA cement (also belite-rich) manufacturing is capable of reducing CO\textsubscript{2} emission and energy consumption by 30% and 20%, respectively\textsuperscript{5}.

The main binding phase of hardened PC pastes and concretes is a calcium silicate hydrate (C-S-H) gel. The structure and properties of C-S-H govern the mechanical properties and durability of PC-based materials\textsuperscript{6,7}. The well-accepted molecular models of C-S-H (Figure 1) are created by modifying a tobermorite structure\textsuperscript{8} with defected dreierketten-type silicate chains, which are composed of bridging and pairing tetrahedra, flanked on CaO\textsubscript{7} sheets to form the calcium silicate basal layer. The neighboring basal layers are separated by zeolitic calcium ions and water\textsuperscript{9}. The structures of synthetic nanocrystalline C-S-Hs have been extensively studied at the atomic, molecular, and nano scales. However, the atomic and molecular structures (e.g., Ca and Si coordination) of hydration product C-S-H have been inadequately studied due to their poor ordering along the c-axis. Understanding the structure and properties of C-S-H is essential for optimizing the properties of sustainable cement-based materials.
Figure 1. Schematic representation of the nanostructures of C-S-H. The Brown circles represent Ca species in the Ca-O sheet, and blue triangles are silicate units in paired and bridging sites. The grey diamonds and yellow squares represent sites which can be occupied by cations that charge-neutralize the structure as a whole (e.g., Ca$^{2+}$ and/or K$^+$).

The hydration of C$_2$S is similar to the intensively studied hydration of C$_3$S; both yield C-S-H gel and portlandite (Ca(OH)$_2$) as hydration products$^{10}$. β-C$_2$S is the most common polymorph of C$_2$S in cements. Using the original surface of the unreacted particle as a reference, C-S-H exhibits a distinct morphology in the outer and inner regions. The inner product (Ip) C-S-H from β-C$_2$S hydration is spherulite-like on the nano-scale$^{11}$, while the outer product (Op) C-S-H is fibrillar$^{12}$ and contributes to the binding properties. Apart from the apparent morphological
difference due to the available space for stacking \(^\text{11}\), there is no significant difference in composition between the Ip and Op \(^\text{12}\). The reactivity of \(\beta\)-C\(_2\)S is lower than C\(_3\)S, but the \(\beta\)-C\(_2\)S hydration is significant for the long-term performance of PC concrete \(^\text{13}\). The \(\beta\)-C\(_2\)S hydration has been investigated by nuclear magnetic resonance (NMR) \(^\text{14}\) \(^\text{15}\) \(^\text{16}\) \(^\text{17}\), electron microscopy \(^\text{18}\) \(^\text{19}\) \(^\text{20}\), and other spectroscopies \(^\text{21}\) \(^\text{22}\) \(^\text{23}\) \(^\text{24}\). However, these techniques are not capable of locally probing the polymerization degree of the silicate chain of the Ip or Op, e.g., NMR, Raman spectroscopy, and infrared spectroscopy can provide chemical environmental Si of C-S-H, but are not capable of providing adequate spatial data \(^\text{25}\) \(^\text{26}\) \(^\text{27}\). Scanning electron microscopy provides high-resolution images, whereas the polymerization degree of C-S-H cannot be evaluated \(^\text{28}\). The potential artifacts generated by the harsh vacuum and high energy electron beam are also not completely understood \(^\text{29}\) \(^\text{30}\) \(^\text{31}\). Recently, \(\alpha\)'\(-\)H-C\(_2\)S, another polymorph of C\(_2\)S, has increasingly gained research and industrial interests due to its higher hydration reactivity compared to \(\beta\)-C\(_2\)S \(^\text{32}\). However, the current understanding of the micro-chemistry (e.g., local coordination environment and silicate polymerization) during its hydration is limited \(^\text{33}\). Understanding the hydration of C\(_2\)S polymorphs and the structure of their hydration products C-S-H is essential for optimizing the properties of sustainable cement and concrete and for lowering the CO\(_2\) emission and energy use in the construction industry.

Scanning transmission X-ray microscopy (STXM) is a powerful technique for characterizing the micro-chemistry of composite materials with a heterogeneous microstructure. It simultaneously provides morphological information and X-ray absorption near-edge structure (XANES) spectra of the elements of interest \(^\text{34}\) \(^\text{35}\), with a spatial resolution of up to 10 nm and a spectral resolution of 0.1 eV \(^\text{36}\). C-S-H gel is vulnerable to electron beam damage \(^\text{37}\), and the soft X-ray used in STXM (< 2000 eV) is more likely to preserve the morphology of C-S-H \(^\text{38}\). The application of
STXM in cement research started in recent years; however, no STXM or XANES study has been reported on the hydration of C$_2$S.

In this work, the hydration products of C$_2$S polymorphs (β- and α’$_H$-) were studied by STXM at the Ca $L_{2,3}$- and Si $K$-edges. The coordination of Ca of anhydrous C$_2$S and C-S-H was also analyzed. Silicate polymerization of the Op and Ip and their C$_2$S interfaces at two different ages were locally probed. Ptychographic imaging was used to provide the interface morphology between the Ip and Op and between water and the Op with a spatial resolution of 10 nm.

**MATERIALS AND METHODS**

Preparation of β- and α’$_H$-C$_2$S. Chemical analytical grade CaCO$_3$, SiO$_2$, and K$_2$CO$_3$ were mixed according to the stoichiometric composition of 1.80CaO-0.10K$_2$O-SiO$_2$ to prepare monoclinic β-C$_2$S. The mixed powder was calcined at 1500 °C for 3 hours and then cooled in air. Chemical analytical grade Ca$_2$SiO$_4$ and Ca$_3$(PO$_4$)$_2$ were stoichiometrically mixed as 1.946CaO-0.892SiO$_2$-0.054P$_2$O$_5$ to prepare α’$_H$-C$_2$S. The mixed powder was sintered at 1450 °C for 6 hours, followed by air cooling. Note that, to ensure the fast formation of single phase (high purity) of β- and α’$_H$- C$_2$S, the calcination temperatures of this laboratory synthesis were higher than typical calcination temperature of belite-rich cement (e.g., 1250 °C). The C$_2$S pellets were then grounded into powders. The particle size distribution D-values, D$_{10}$, D$_{50}$, and D$_{90}$ of anhydrous β-C$_2$S were 1.19 μm, 6.74 μm, and 18.48 μm, respectively. The D$_{10}$, D$_{50}$, and D$_{90}$ of anhydrous α’$_H$-C$_2$S were 1.45 μm, 7.05 μm, and 19.08 μm, respectively.
Sample preparation for STXM. Anhydrous C$_2$S powders were mixed with deionized water with a water-to-solid ratio (w/s) of 10 so that the particles were easily isolated by drop-casting to satisfy the transmission observation in the STXM. The suspensions were stored in polyethylene tubes filled with N$_2$ gas at 25±1 °C. At early and late curing ages (10 days and 40 days for α’-H-C$_2$S; 17 days and 51 days for β-C$_2$S), the suspension of 0.5 μL was dropped onto a Si wafer frame (5 mm × 5 mm, 100 nm thick), where a 100-nm-thick Si$_3$N$_4$ membrane window (1 mm × 1 mm, 100 nm thick) is in the center. The liquid was quickly and gently absorbed with a tip of KimWipes™ from the edge of the Si frame. The liquid removal process was monitored under an optical microscope to ensure that most of the solids remain on the membrane window. The anhydrous powders were dry-dusted on the Si$_3$N$_4$ membrane windows directly. These windows were then placed on the STXM sample stage inside a vacuum chamber.

Scanning Transmission X-ray Microscopy. The XANES spectra and images at Si K-edge (1830–1880 eV) and Ca L$_{2,3}$-edge (342–360 eV) were obtained at beamlines 5.3.2.1 and 5.3.2.2, respectively, at the Advanced Light Source of the Lawrence Berkeley National Laboratory (LBNL). Comprehensive descriptions of the STXM beamlines are given in 36 48. The optical density (OD) at a specific energy of all pixels is affected by composition, density, and sample thickness 40. Element mappings were obtained by differential measurements of the OD at pre- and on- edge at a spatial resolution up to 25 nm. Image stacks were collected by scanning the region of interest (ROI) at a series of beam energies, which yielded a XANES spectrum of each pixel on the ROI. RGB overlay maps, which visualize multiple phases with
various compositions, were generated using singular value decomposition with reference spectra. All data processing was conducted using aXis2000 software. At each age, at least two RGB maps were taken to obtain statistically reliable results, and at least ten spectroscopic measurements were taken from different particles on each Si₃N₄ membrane.

Ptychographic images were collected at 800 eV at beamline 5.3.2.1. Full details of the imaging technique are given in. The spatial resolution of these transmission images is ~10 nm. The images were reconstructed by Nanosurveyor package. The width of C-S-H fibers of hydrated samples in the ptychographic images was quantified using ImageJ with F3D plugin developed at LBNL.

RESULTS AND DISCUSSION

Ca coordination of anhydrous β-C₄S and hydrated β-C₄S for 17 days. The Ca L₂,₃-edge XANES spectra (Figure 2) consist of two major peaks, L₃ 2P₃/2 (a₂) and L₂ 2P₁/2 (b₂), which are due to the loss of degeneracy of 2p orbitals by spin-orbital interactions, along with a number of minor leading peaks, a₀, a₁, and b₁, in order of increasing energy. The peak energy difference of a₂-a₁ and b₂-b₁ (ΔL₃ and ΔL₂, termed as splitting energy) and the intensity of the minor peaks are relevant to the coordination symmetry of Ca in the first shell and its ordering.
Figure 2. Hydrated $\beta$-C$_2$S at 17 days: (A) Transmission image of the ROI taken at 349 eV; (B) selected regions in the Ca element map for the XANES spectra; (C) Ca $L_{2,3}$-edge XANES spectra extracted from different regions specified in (B).

The Ca $L_{2,3}$-edge XANES spectrum (Figure 2C) of anhydrous $\beta$-C$_2$S (monoclinic) exhibits a cubic-like symmetry with peak $a_{10}$ located between $a_1$ and $a_2$, and the peak energy differences of $a_2-a_1$ and $b_2-b_1$ are 1.54 and 1.2 eV, respectively. $\beta$-C$_2$S contains seven-fold and eight-fold coordinated Ca, denoted as Ca1 and Ca2 (Supporting Information, Figure S1). The equatorial plane of Ca1 is parallel to (10\overline{1}) with Ca-O bond lengths for Ca1 ranging from 2.273 to 2.892 Å with an average of 2.528 Å. The five equatorial O atoms are not exactly coplanar. The Ca2 is
coordinated by eight O atoms with a distorted symmetry for the cube, where the two trapezoids formed of the top four and bottom four O atoms are misaligned along each other’s diagonal direction by 55°. The Ca-O bond lengths for Ca2 range from 2.400 to 2.676 Å with an average of 2.503 Å. The complex coordination of Ca1 and Ca2 in β-C2S produces weak cubic-like crystal field splitting effects.

To avoid X-ray absorption saturation at low energy, only particles with size of ~1 μm (Figure 2A and 1B) were measured at Ca L2,3-edge. The 17-day hydrated β-C2S grains at the Ca L2,3-edge show mostly identical peak positions and an equivalent splitting energy in each grain, indicating that the Ca in each grain exhibits a similar environment. Peaks a2 and b2 in the selected areas are 0.1 eV lower than anhydrous β-C2S, suggesting a lower bond strength of Ca-O. The splitting energies in the selected areas are 1.1 eV, suggesting a distorted octahedral-like symmetry for Ca52. The X-ray absorption features (i.e., splitting energy and peak positions) are similar to 14Å tobermorite (calcium silicate hydrate mineral, [Ca4Si6O16(OH)2 · 2H2O]·(Ca·5H2O))57 58, suggesting the C-S-H formed in this hydration system is structurally analogous to tobermorite. The relative intensity ratios of peak a1-to-a2 (I(a1)/I(a2)) and b1-to-b2 (I(b1)/I(b2)) of these selected areas are 0.021-0.041 and 0.050-0.077, respectively, which are lower than hydrothermally synthetic C-S-H at 25 °C (0.05-0.15 for I(a1)/I(a2) and 0.07-0.15 for I(b1)/I(b2)) 38. The details in the calculation of relative intensity of these peaks are shown in Figure S2 (Supporting information). Synthetic C-S-H has a longer range order than C-S-H formed in the β-C2S hydration along the c-axis 59. This suggests a shorter range order for Ca in this hydration system since the Ca/Si molar ratio of C-S-H in hydrated β-C2S ranges from 1.2 to 2.1 60. Moreover, no contribution by portlandite (a non-strength-giving phase in hydrated C2S) was observed in the spectra, suggesting that C-S-H does not intermix with portlandite at the 25 nm-scale in this ROI,
and the interlayer six-fold coordinated Ca is also distorted. Thus, the Ca of C-S-H in hydrated β-C₂S is short-range-ordered and tobermorite-like. Portlandite crystallites were observed elsewhere in these dilute systems (Supporting information, Figure S3).

Hydrated C₃S at 17 days also exhibited comparable peak positions, splitting energies, and intensity ratios in XANES at the Ca L₂,₃-edge, indicating that the Ca environment of hydrated C₃S is similar to hydrated β-C₂S. The extended X-ray absorption fine structure study showed that the Ca-O environment of hydrated β-C₂S is similar to 11Å tobermorite, but the coordination number and Ca-O bond length of C-S-H could not be determined due to the presence of portlandite. An ¹⁷O NMR study demonstrated that Ca-OH sites are present in C-S-H of hydrated β-C₂S. The C-S-H structure of hydrated β-C₂S proposed by Richardson contains six- and seven-fold coordinated Ca in the interlayer and intralayer, respectively. The model proposed by Gartner suggests that the interlayer Ca of C-S-H from hydration of C₃S and β-C₂S is six-fold coordinated. In the present work, the distorted six-fold Ca in the interlayer of hydrated β-C₂S fits both models.

Ca coordination of anhydrous α’-H-C₂S and α’-H-C₂S hydrated for 10 days. The Ca L₂,₃-edge XANES spectra of α’-H-C₂S (orthorhombic) also exhibits a cubic-like symmetry with peak energy differences of a₂-a₁ of 1.57 eV and b₂-b₁ of 1.25 eV in Figure 3. This indicates a slightly stronger crystal field of Ca for α’-H-C₂S. α’-H-C₂S contains nine- and eight-fold coordinated Ca at two sites (denoted as Ca1 and Ca2, see Figure S1 in Supporting information) with Ca-O bond lengths for Ca1 ranging from 2.214 to 3.175 Å with an average of 2.708 Å, and those for Ca2 ranging from 2.292 to 2.927 Å with an average of 2.545 Å. The superposition of symmetries and complex
coordination leads to the cubic-like crystal field splitting of $\alpha'_H$-$C_2S$. The energy difference in $\alpha'_H$-$C_2S$ is slightly larger compared to $\beta$-$C_2S$ (monoclinic), suggesting that Ca-O complexes in $\alpha'_H$-$C_2S$ are more similar to cubic-like symmetry (less distorted) than in $\beta$-$C_2S$. For Ca1 in $\beta$-$C_2S$, its coordination number is seven, which is incapable of providing an ideal cubic symmetry.

Figure 3. Hydrated $\alpha'_H$-$C_2S$ at 10 days: (A) Transmission image of the ROI taken at 349 eV; (b) selected regions in the Ca element map for the XANES spectra; (c) Ca $L_{2,3}$-edge XANES spectra extracted from different regions specified in (b).

Portlandite was not observed in this region for $\alpha'_H$-$C_2S$ hydrated for 10 days. Mostly identical peak positions and peak intensities were observed in each grain. Peaks a$_2$ and b$_2$ in the selected areas are ~ 0.1 eV lower than for $\alpha'_H$-$C_2S$, and the peak positions are 0.05 eV higher than for
hydrated β-C₂S. The splitting energies in the compounds are ~0.1 eV lower than for α’-H-C₂S but
~0.05 eV higher than for hydrated β-C₂S. This suggests that the bond strength of Ca-O of
hydrated α’-H-C₂S is slightly higher than for hydrated β-C₂S; namely, more six-fold coordinated
Ca is present in the α’-H-C₂S hydration system. The intensity ratios of hydrated α’-H-C₂S are
0.018-0.038 and 0.037-0.067, which are also lower than for anhydrous but comparable to or
slightly lower than hydrated β-C₂S. These absorption features indicate that the coordination
symmetry of Ca in hydrated α’-H-C₂S is also tobermorite-like, but the ordering of Ca is slightly
lower than for hydrated β-C₂S.

**Si environment of anhydrous β-C₂S and β-C₂S hydrated for 17 days.** The major peak of a₁ at
the Si K-edge of β-C₂S (Figure 4C) is assigned to the electronic excitation from 1s to the
antibonding 3p-like state orbital (t₂) of tetrahedral Si⁶⁴⁶⁵. Pre-edge peak a₀ is attributed to the
dipole-forbidden transition of 1s electrons to the antibonding 3s-like states⁶⁶. This forbidden
peak appears due to the mixing of s- and p- states when the SiO₄ tetrahedra coordination is
highly distorted. Thus, the intense a₀ peak of β-C₂S suggests a strong distortion of silicate
tetrahedron. The minor peak of a₂ is assigned to the multiple-scattering effect from more distant
atom shells through a photoelectron interaction; its energy is governed by the interatomic
distance and structural complexity⁶⁷.

The interface between the β-C₂S core and the hydrous layer C-S-H is well distinguished (Figure
4), indicating changes in the phase density. The OD of the hydration product layer slightly
decreases with increasing distance from the interface, which is associated with several small
brighter regions. Compared to C₃S hydration system⁶¹, the C-S-H of hydrated β-C₂S is more
dispersed, and fibrils are not present at this age and at this resolution (Figure 4A).

Figure 4. Hydrated $\beta$-C$_2$S at 17 days: (A) Transmission image of the ROI taken at 1849 eV; (B)
selected regions in the Si element map from the XANES spectra; (C) Si K-edge XANES spectra extracted from different regions specified in (B); (D) RGB overlay map using XANES spectra obtained from the C-S-H (Area 4, green) marked in (B) and reference spectrum of anhydrous β-C₂S (red). More RGB over maps can be found in Figure S4 (Supporting information).

Peak a₁ is not shown at ~1851.1 eV in areas 3, 4, 6 but a single a₁ peak is at 1848.8 eV in these areas (Figure 4C), suggesting that C-S-H is the only silicate phase in these areas. The absorption features of C-S-H in hydrated β-C₂S resemble those of C-S-H in C₃S hydration systems ⁶⁸ ⁶¹ ⁴³. The Si K-edge (a₁ peak) at 1848.8 eV of C-S-H is 2.2 eV lower than that of β-C₂S due to the longer Si-O bond of the C-S-H relative to β-C₂S ⁶⁹ ⁵⁶. Li ⁶⁴ concluded that although there is considerable overlapping for silicate minerals with different polymerization types, the Si K-edge generally shifts to higher energy with increased silicate polymerizations. Thus, in this case, the Si-O bond length, rather than the polymerization degree, dominates the Si K-edge shift.

The energy difference between a₁ and a₂, Δa₂-a₁, can be used to evaluate the degree of silicate polymerization ³⁸,⁴³,⁶¹,⁶⁸. The values of Δa₂-a₁ for areas 3 and 4 are equivalent, suggesting that the degree of polymerization of the C-S-H here is comparable. The energy positions of multiple scattering peaks (e.g., a₂) are affected by the Si interatomic distances, while the intensity and number of resonances are heavily dependent on the bond angle, site symmetry, and number of nearest neighboring oxygen atoms (Si in most cement phases are 4-fold coordinated, expect thaumasite, where Si is 6-fold coordinated) ⁷⁰. Two a₂ peaks are distinguished in the spectra in areas 2 and 5. In these areas, a peak at 1848.8 eV and a broad shoulder/hump at ~1851.1 eV were observed, the peak is assigned to the a₁ peak of C-S-H, and the shoulder/hump is the contribution of a₁ peak (at 1851.0 eV) of anhydrous β-C₂S. These absorption features suggest that areas 2 and 5 are mixtures of C-S-H and anhydrous β-C₂S, and the hydration here is incomplete. Similarly,
areas 7 and 8 are mixes of C-S-H and anhydrous \(\beta\)-C\(_2\)S, implying that small \(\beta\)-C\(_2\)S particles exist in these areas. This fact is consistent with the original granulometry result of \(\beta\)-C\(_2\)S, \(~10\%\) of \(\beta\)-C\(_2\)S was below 1 \(\mu\)m. Note that a weak contribution of C-S-H to the spectrum at \(~1848.8\) eV is observed in area 1 since C-S-H surrounds the entire grain. \(\Delta a_{2-a_1}\) of area 6 is 0.9 eV lower than those of areas 3 and 4 (\(~15.7\) eV), suggesting a lower degree of polymerization of C-S-H in area. The difference in local polymerization degree can be explained by the different dissolution rates on the crystal faces and by different numbers of surface defects of \(\beta\)-C\(_2\)S \(^{71}\); multiple crystal faces with low hydraulic reactivity occur in area 6. Note that, hydraulic reactivity of all faces of pure \(\beta\)-Ca\(_2\)SiO\(_4\) is similar; the dopant changes the local reactivity of \(\beta\)-C\(_2\)S crystals \(^{72}\). Similarly, the value of \(\Delta a_{2-a_1}\) for C-S-H in area 9 is 0.7 eV lower than for C-S-H in the unconstrained space. The RGB overlay map (Figure 4d) shows that the C-S-H surrounds the unreacted core, which is consistent with the analysis above. The values of \(\Delta a_{2-a_1}\) for C-S-H among different grains (Areas A- D) are 15.6-16.1 eV, indicating a distinction in silicate polymerization due to the interference of other grains and the hydration rate. In addition, the contribution of cross-linking sites to the spectra is not observed in the typical range at 1853-1854 eV \(^{73}\) since the basal layers of C-S-H are separated by \([\text{Ca(H}_2\text{O)}_6]\)^{2+} or \([\text{Ca(OH)(H}_2\text{O)}_5]\)^+ \(^{63}\). \(^{29}\)Si NMR studies showed that the mean silicate chain lengths of C-S-H of 7-12 months hydrated \(\beta\)-C\(_2\)S are still lower than 3.5 \(^{12}\) \(^{74}\). This suggests that the proportion of long silicate chains (octamer or longer) in this C-S-H is relatively low. Therefore, the C-S-H at 17 days is mainly a mixture of dimer and pentamer C-S-Hs.

It is well known that the degree of silicate polymerization of synthetic C-S-H increases as Ca/Si decreases \(^{27}\) \(^{26}\). Although the degree of polymerization of hydrated \(\beta\)-C\(_2\)S progressively increases over time \(^{17}\) \(^{75}\), there is no significant variation in the mean Ca/Si of Op (excluding intermixed
Portlandite) with age \(^{37}\). Thus, it is difficult to estimate the local degree of polymerization based on Ca/Si ratio. STXM is a more reliable technique to locally probe the variation in silicate polymerization of the C-S-H without considering intermixed microcrystalline portlandite in the hydrous regions \(^{12}\) and unreacted \(\beta\)-C\(_2\)S.

**Si environment of hydrated \(\beta\)-C\(_2\)S for 51 days.** Two distinct optical densities were observed at 51 days (Figure 5A). Double \(a_1\) peaks are not observed in Figure 5B, and only a single \(a_1\) peak at 1848.9-1849.1 eV occurs. Peak \(a_0\) at 1847.9 eV was not observed, suggesting a fully hydrated grain (see Supporting information, Figure S5). The peak difference of \(\Delta a_2-a_1\) for areas 5, 6, 7, 8, 9, and B is 16.7-17.0 eV, and 15.8-16.2 eV for areas 1, 2, 3, and 4; two distinct ranges of \(\Delta a_2-a_1\) were observed in Figure 5C. Thus, the two distinct ODs indicate the existence of Ip (dark region) and Op (light region) in Figure 5A. Areas 5, 6, 7, 8, and 9 are Ip dominant, while the areas 1, 2, 3, and 4 are Op dominant. The Si \(K\)-edge \((a_1)\) and \(\Delta a_2-a_1\) of Ip are both lower compared to those of Op, indicating a lower silicate polymerization and a slightly longer Si-O bond for Ip. The \(\Delta a_2-a_1\) and Si \(K\)-edge for Op at 51 days are slightly higher than at 17 days, revealing that the Op polymerizes over time. The lower \(\Delta a_2-a_1\) for areas 10 and A may be explained by impingement from adjacent grains. Similarly, contribution of cross-linking sites to the spectra was also not observed due to the abundant interlayer Ca between basal layers of C-S-H \(^{63}\). Thus, both Op and Ip are likely mixtures of dimer and pentamer C-S-Hs with very small amounts of octamer and longer chains. The values of \(\Delta a_2-a_1\) for Op and Ip are larger compared to those of synthetic C-S-H with a bulk Ca/Si molar ratio of 1.4 \(^{38}\), suggesting that the electron shielding on Si from cations in hydration product C-S-Hs is weaker. This can be explained by the highly distorted Ca of \([\text{Ca(OH)}(\text{H}_2\text{O})_5]^+\) in the interlayer of C-S-Hs; and the partial replacement of divalent Ca by
monovalent K from the dopant\textsuperscript{76}. Future study will focus the coordination environment of K in this system and the electron density around these atoms.

Figure 5. Hydrated \(\beta\)-C\textsubscript{2}S at 51 days: (A) Transmission image of the ROI at 1849 eV; (B) selected regions in the Si element map for the XANES spectra; (C) Si K-edge XANES spectra
extracted from different regions specified in (C); (D) ptychographic image of another area at the same age. Op is indicated by blue arrows, Ip is indicated by red arrows, and portlandite is indicated by yellow arrows.

The ptychographic image (Figure 5d) shows the fibrillar Op of another hydrated β-C₂S particle at 51 days; the width of fibers has a distribution of 11-25 nm, which is smaller than the Op of C₃S. The Ip of β-C₂S exhibits a dense aggregated structure, and the Ip of C₃S with the same w/s exhibits a less dense structure with aggregates of globules. The morphological difference between the two Ips can be explained by the lower reactivity of β-C₂S. Geng reported that the Ip and Op of hydrated C₃S exhibits an identical Si K-edge with a higher energy for the a₂ peaks of Op, showing that Ip has a lower degree of silicate polymerization. Portlandite microcrystallites intermixed with Op is observed, which is consistent with TEM study of Richardson.

**Si environment of anhydrous and hydrated α’₇-H-C₂S for 10 days.** Similar to β-C₂S, pre-edge peak a₀ was also observed in α’₇-H-C₂S (Figure 6c), suggesting a slight distortion of the silicate tetrahedron of α’₇-H-C₂S. The Si K-edge of α’₇-H-C₂S is 0.2 eV higher than for β-C₂S, suggesting a slightly shorter Si-O bond for α’₇-H-C₂S compared with β-C₂S, which is consistent with XRD refinement.
Figure 6. Hydrated $\alpha'_H$-C$_2$S at 10 days: (A) Transmission image of the ROI taken at 1849 eV; (B) selected regions in the Si element map for the XANES spectra; (C) Si K-edge XANES spectra extracted from different regions specified in (B); (D) RGB overlay map using XANES spectra obtained from the C-S-H (area 7, green) marked in (B) and reference spectrum of
anhydrous $\alpha_{\text{H}}'$-C$_2$S (red).

Similar to $\beta$-C$_2$S hydration, the interface between non-fibrillar Op and $\alpha_{\text{H}}'$-C$_2$S was observed at 10 days (Figure 6). The Si $K$-edge and $\Delta a_2$-$a_1$ of hydrated $\alpha_{\text{H}}'$-C$_2$S at 10 days are equivalent to or 0.2-0.3 eV larger than for hydrated $\beta$-C$_2$S at 17 days. Thus, the chemical environment of Si in hydrated $\alpha_{\text{H}}'$-C$_2$S is similar to that in $\beta$-C$_2$S hydration at 17 days with a comparable degree of polymerization at an earlier age. The polymerization degree of the Op is more homogeneous since the $\Delta a_2$-$a_1$ values of all C-S-Hs here are ~15.9 eV. Again, Op cross-linking sites are not present, suggesting that the Op is still mainly a mixture of dimer and pentamer units. Thus, $\alpha_{\text{H}}'$-C$_2$S exhibits a higher reactivity relative to $\beta$-C$_2$S. This work is consistent with experimental and simulation studies on the reactivity but provides new details on the rate of polymerization of $\alpha_{\text{H}}'$-C$_2$S hydration.

**Si environment of hydrated $\alpha_{\text{n}}'$-C$_2$S for 40 days.** The C-S-H of $\alpha_{\text{H}}'$-C$_2$S at 40 days is fibrillar in Figure 7. There was no Ip/Op interface in the fully hydrated grain (see Supporting information, Figure S6) observed. The values of $\Delta a_2$-$a_1$ for all the areas range from 16.7 to 17.0 eV, suggesting a relatively homogenous polymerization. This is a primary difference compared to $\beta$-C$_2$S. The Si $K$-edge ($a_1$) of all areas is comparable, suggesting similar Si coordination in these areas. The comparable degree of polymerization of the interfaces between different grains suggests that the polymerization is not obviously interfered by adjacent grains. Additionally, the C-S-Hs are mainly mixtures of dimer and pentamer units, and a cross-linking effect was not observed on the spectra. Compared to the Op of $\beta$-C$_2$S at 51 days, hydrated $\alpha_{\text{H}}'$-C$_2$S at 40 days
exhibited a comparable polymerization degree at an earlier age. This confirms the higher reactivity of $\alpha'_H$-C$_2$S compared with $\beta$-C$_2$S.

Figure 7. Hydrated $\alpha'_H$-C$_2$S at 40 days: (A) Transmission image of the ROI taken at 1849 eV; (B) selected regions in the Si element map for the XANES spectra; (C) Si K-edge XANES spectra extracted from different regions specified in (B); (D) ptychographic image.

The ptychographic image (Figure 7D) shows the fibrillar C-S-H with a width of 15-30 nm, which
is larger than hydrated $\beta$-C$_2$S but smaller than hydrated C$_3$S (41-59 nm)\(^{77}\), and the inner C-S-H exhibits a looser aggregated structure with $\sim$17 nm voids compared to the Ip of hydrated $\beta$-C$_2$S. The Ips of hydrated $\beta$-C$_2$S, C$_3$S, and PC with a practical w/s (i.e., <0.6) appear to be small and globular\(^{79}\). The larger void of the Ip relative to the Ip of hydrated C$_3$S\(^{79}\) could be induced by the w/s of 10 used in this study, which provides sufficient space for nanocrystallite growth during the fast hydration. The microstructure of hydrated $\alpha'$$_H$-C$_2$S is similar to hydrated C$_3$S with the same w/s\(^{77}\) instead of the hydrated $\beta$-C$_2$S mentioned above. Thus, the morphological difference among the silicates in these diluted systems may be relevant to their hydration reactivity. More ptychographic images can be found in Figure S7 (Supporting information).

**CONCLUSIONS**

The coordination environments of Ca and Si in anhydrous $\beta$-C$_2$S, $\alpha'$$_H$-C$_2$S, and their hydration products were studied using STXM; the morphology of C-S-Hs was imaged by ptychography. The key conclusions are:

1. Ca in both anhydrous phases are in a distorted cubic symmetric coordination. The Ca coordination of C-S-H in both hydration systems is highly distorted and is similar to 14Å tobermorite but have a shorter-range order. More six-fold coordinated Ca occur in hydrated $\alpha'$$_H$-C$_2$S than in hydrated $\beta$-C$_2$S.

2. The Si of both anhydrous phases have asymmetric tetrahedral coordination. The silicate chain of the Op in both hydration systems polymerizes over time. The hydration products in the $\alpha'$$_H$-C$_2$S system polymerize faster than in $\beta$-C$_2$S. The hydraulic reactivity of $\alpha'$$_H$-C$_2$S is higher.
(3) Ip has a lower degree of silicate polymerization than Op in fully hydrated $\beta$-C$_2$S, which is accompanied with a slightly longer Si-O bond due to the space constraint for contacting water. Compared with hydrated $\beta$-C$_2$S, the silicate polymerization in hydrated $\alpha'_H$-C$_2$S is more homogenous.

(4) The C-S-H of hydrated $\beta$-C$_2$S exhibits fine fibrils that intermix with portlandite, and Ip appears to be dense aggregated structure. The morphology of fully hydrated $\alpha'_H$-C$_2$S is more homogenous, and its inner region is slightly more porous with nano-size voids. Ip and Op are not distinguished in fully hydrated $\alpha'_H$-C$_2$S

Therefore, this study provides insights into advanced applications of the hydration behaviors of C$_2$S to C$_2$S rich cements and the controllability of the performance of these materials in service. The lower calcination temperature and CO$_2$ emission of $\alpha'_H$-C$_2$S during manufacturing can reduce the environmental impact of cement manufacturing. Due to its high reactivity, $\alpha'_H$-C$_2$S can be a practical alternative to C$_3$S and $\beta$-C$_2$S in Portland cement or BCSA cement. Therefore, this work is an important step towards the development of hydration mechanism of C$_2$S-rich cement, which will provide new insight into the performance of these sustainable cements (lower CO$_2$ emission and energy use) in service.

ASSOCIATED CONTENT

Supporting Information

Coordination environment of Ca in two C$_2$S polymorphs; an example of fitting of a XANES spectrum at Ca $L_{2,3}$-edge; transmission images at 349 eV of samples at 10 and 17 days; RGB
overlay maps of samples at 10, 17, 40, and 51 days; ptychographic images of samples at 40 and 51 days.

AUTHOR INFORMATION

Author Contributions

J.L., G.G., and P.J.M. conceived and designed the experiments. J.L. conducted the experiments. Y.-S.Y. and D.A.S. supported the imaging experiments. J.L., G.G., and W.Z. analyzed the results. J.L. and G.G. wrote the manuscript. All authors reviewed the manuscript.

ACKNOWLEDGEMENTS

This research was founded by the Republic of Singapore’s National Research Foundation through a grant to the Berkeley Education Alliance for Research in Singapore (BEARS) for the Singapore-Berkeley Building Efficiency and Sustainability in the Tropics (SinBerBEST) Program. The Advanced Light Source is supported by the Director, Office of Science, and Office of Basic Energy Sciences of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231. G. Geng also wants to acknowledge European Union’s Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie grant agreement No 701647. We thank Matthew Marcus for insightful discussions on fitting and interpretation of Ca $L_{2,3}$-edge XANES spectra. We thank Ellis Gartner for the personal communication and help on the study of dicalcium silicates.

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SYNOPSIS

The atomic-scale and nano-scale structure of hydrated dicalcium silicate are studied, which may inspire the manufacture of cement with significantly lower CO₂ emission.

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