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**Rachel Cook** 

Hongyan Ma Missouri University of Science and Technology, mahon@mst.edu

Aditya Kumar *Missouri University of Science and Technology*, kumarad@mst.edu

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**Research Article** 

# Mechanism of tricalcium silicate hydration in the presence of polycarboxylate polymers



Rachel Cook<sup>1</sup> · Hongyan Ma<sup>2</sup> · Aditya Kumar<sup>1</sup>

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#### Abstract

The early-age hydration of cement is inhibited in the presence of comb-shaped polycarboxylate ether (PCE) polymer—a dispersant commonly added to control rheological properties of fresh cement paste. This study employs a series of micro-calorimetry experiments and phase boundary nucleation and growth simulations to elucidate the effects of dosage and molecular architecture of PCE on hydration of tricalcium silicate ( $Ca_3SiO_5$  or  $C_3S$  in cement notation), the dominant phase in cement. Results show that PCE—regardless of its molecular architecture—suppresses early-age hydration of  $C_3S$ . PCE-induced retardation becomes increasingly more pronounced as dosage of PCE in the paste increases. Such suppression of  $C_3S$  hydration has been attributed to adsorption of PCE molecules on silicate surfaces, which inhibit topographical sites of  $C_3S$  dissolution and C–S–H nucleation, and impede the post-nucleation growth of C–S–H. This study develops a correlation between molecular architecture of PCE and its ability to suppress  $C_3S$  hydration through quantitative analyses of retardation effects induced by PCEs with different molecular architectures. The numerical equation, describing such correlation, offers a reliable, and, more importantly, a readily quantifiable indicator of PCE's potential to suppress  $C_3S$  hydration in relation to its dosage and molecular architecture. In the context of practical application of this study, the aforementioned numerical equation can be used to order and rank PCEs—of various molecular architectures—on the bases of their potentials to suppress  $C_3S$  hydration, and to select ones that cause the optimum (i.e., user-desired) extent of hydration suppression.

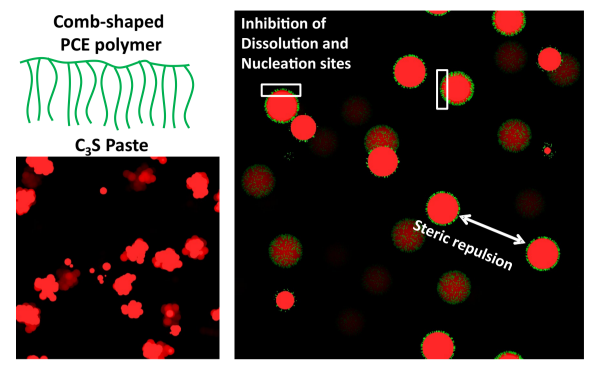
Aditya Kumar, kumarad@mst.edu; Rachel Cook, recwx7@mst.edu; Hongyan Ma, mahon@mst.edu | <sup>1</sup>Department of Materials Science and Engineering, Missouri University of Science and Technology (S&T), B49 McNutt Hall, 1400 N. Bishop, Rolla, MO 65409-0340, USA. <sup>2</sup>Department of Civil, Architectural and Environmental Engineering, Missouri University of Science and Technology (S&T), Rolla, MO 65409-0340, USA.



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#### **Graphical abstract**



**Keywords**  $C_3S \cdot PCE \cdot Hydration kinetics \cdot Nucleation and growth \cdot Simulation \cdot Microcalorimetry$ 

# **1** Introduction

The reaction of cement with water, that is, hydration, involves the occurrence of two concurrent processesdissolution of anhydrous phases present in cement, and precipitation of hydration products (subsequently referred to as hydrates) [1, 2]. In a typical cement paste (i.e., [cement + water] system), the hydrate that occupies the largest volume fraction—and, thus, considered the main hydrate—is calcium-silicate-hydrate (C-S-H, wherein  $C = CaO, H = H_2O$ , and  $S = SiO_2$  as per standard cement notation) [1, 2]. The strong electrostatic bonding between the nanometer-scale components of C-S-H binds the paste cohesively, and lends the solid-to-solid phase connectivity within the paste's microstructure needed for setting and hardening (development of mechanical properties, e.g., compressive strength) [3-6]. In cement pastes, the nucleation of C-S-H occurs in heterogeneous manner on solid substrate boundaries, that is, cement particles' surfaces; as such, the mechanism of its precipitation is typically designated as phase boundary nucleation and growth (pBNG) [3, 7–11]. As properties of cement paste are largely dictated by rate and amount of C–S–H precipitation, factors that affect C-S-H precipitation inevitably affect the

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development of the paste's mechanical properties. One such factor—that imparts significant effect on C–S–H's nucleation and growth—is the presence of polymer-based chemical admixtures in the paste.

Polycarboxylate ether (PCE) superplasticizers are a wellknown class of comb-shaped, polymer-based dispersants, typically used to control the rheological properties of fresh cementitious systems [12, 13] (e.g., high-performance concrete). A singular PCE molecule, if isolated, could be characterized as having comb-shaped architecture, consisting of an anionic backbone—usually formulated using polyacrylic acid or polymethacrylic acid—grafted with a number of hydrophilic ethylene oxide side chains [14-22]. When introduced in cement pastes, PCE's negatively charged backbone adsorb onto positively charged cement particles' surfaces through electrostatic interactions. Meanwhile, the side-chains-which protrude into the solution, oriented away from cement particles' surfaces-induce steric hindrance between neighboring cement particles, thus alleviating the effects of particle agglomeration [14-19]. The plasticizing mechanism of PCE also induces a side-effect—retardation of hydration kinetics of cement [14, 15, 23-25]. More specifically, the adsorption of PCE onto cement particles' surfaces inhibits topographical cement dissolution and C-S-H nucleation sites, thus suppressing cement's reactivity as detailed in the literature [15, 16, 18, 24, 25]. There is consensus among researchers that the molecular architecture of

PCE—specifically, the number of side chains grafted onto each unit of backbone (n), the carboxylate-to-ether ratio (C/E), and the number of ethylene oxide monomers constituting the side chain (P)—significantly affects PCE's adsorption behavior, and, thus, its potential to suppress cement hydration [15-17, 20, 22]. Notwithstanding, the exact mechanisms of PCE-cement interactions, especially in relation to PCE's molecular architecture, are not well understood. Several prior studies have argued that the adsorption capacity of PCE is largely dependent on its side chain grafting density (i.e., inverse of C/E: carboxylate-to-ether ratio), wherein lower grafting densities (or higher C/E) entail higher residual negative charge on the backbone, and, thus, improved adsorption onto positivelycharged cement particles' surfaces [15, 26-28]. Other studies, however, have argued that the length of the side chain (i.e., P: number of monomers constituting each side chain)—as opposed to the side chain grafting density has greater influence on the PCE's adsorption capacity [17, 29]. The premise, here, is that shorter side chains ensure that accessibility to the negative charges on the PCE's backbone is not hindered or limited by steric hindrance (induced by its side chains); this enables better adsorption of PCE molecules onto cement particles' surfaces. In contrast to the above, some studies [15, 16, 30] have posited that the PCE's charge density—which acquires higher values at lower side chain grafting densities (or higher C/E) and shorter side chain lengths (P)—influences the PCE's adsorption capacity the most. More specifically, higher charge density leads to stronger, and better-distributed, electrostatic interactions between cement particles and PCE molecules, and, therefore, improved adsorption. In a recent study, Marchon et al. [22] reported that a more encompassing, composite architectural parameter-that accounts for the aforementioned parameters (i.e., C/E, n, and P) as well as the molecular weight and dosage of PCE in the system—is required to fully describe the effects of PCE on C<sub>3</sub>S hydration. The same authors [21] also showed that for a series of cement pastes, provisioned with different dosages of different PCEs, the composite architectural parameter scaled, broadly in a monotonic fashion, with respect to delay in occurrence of the maximum hydration rate. In this study, focus is given to rigorously test the ability of the composite architectural parameter to reliably quantify the retardation caused by PCEs of different molecular architectures.

In addition to aforementioned knowledge gaps pertaining to correlations between PCE's molecular architecture and its ability to suppress cement hydration—the effect of PCE on nucleation and growth of the main hydrate, i.e., C–S–H, is still not well understood. In recent studies [23, 24, 31], it has been suggested that in cementitious paste provisioned with PCE, due to blockage of nucleation sites by PCE molecules, C-S-H nucleation changes from heterogeneous (i.e., on solid surfaces) to homogeneous (i.e., in the pore space). The authors [23, 24] argued that the change necessitates higher supersaturation for C-S-H precipitation, thus enforcing prolonged dissolution of cement until massive precipitation of C-S-H can occur. However, in other studies [16, 30], it has been shown that while PCE changes the nucleation and growth processes' rate constants (e.g., rate of growth, frequency of nucleation sites), the precipitation of C-S-H still occurs in heterogeneous manner. In a more recent study, Meng et al. [25] argued that even at higher PCE dosages, nucleation and subsequent growth of C-S-H continues to occur heterogeneously, albeit at supressed rates. The authors [25] reported that the suppression is caused by: (1) adsorption of PCE molecules onto cement surfaces-which blocks a fraction of C-S-H nucleation sites, and (2) adsorption of PCE molecules onto C-S-H—which partially blocks C–S–H's access to the pore solution, resulting in the inhibition of its post-nucleation growth.

The above discussion highlights the current state of knowledge, as well as gaps in knowledge, pertaining to underlying mechanisms that link PCE's molecular architecture to its ability to suppress cement hydration and nucleation and growth of C-S-H. The main reason, that would explain these knowledge gaps, is that majority of the past studies have examined the role of PCE in multi-component cementitious systems, in which it is infeasible to de-couple the effects of PCE on dissolution-precipitation hydration process of the two most reactive cement clinker mineral phases, that is, tricalcium silicate (C<sub>3</sub>S) and tricalcium aluminate (C<sub>3</sub>A, where A: Al<sub>2</sub>O<sub>3</sub> as per standard cement notation). For example, in such multi-component cement systems, C<sub>3</sub>A hydrates rapidly in the presence of gypsum to form ettringite—which then serves as a favorable adsorbent for PCE molecules [17, 18, 32]. As PCE is drawn in substantial amounts from the solution and adsorbed onto ettringite, the influence of PCE on C<sub>3</sub>S hydration rates is marginalized and, therefore, difficult to isolate from the overall response. Furthermore, in multi-component systems, interactions between PCE-and-C<sub>3</sub>A and PCE-andettringite may affect (i.e., increase or decrease) the amount of free aluminate  $[Al(OH)_{4}^{-}]$  ions in the solution—which, in turn, makes it difficult to isolate and analyze the net effect of PCE (vis-à-vis that of aluminate ions) on C<sub>3</sub>S hydration rates [20, 21]. Therefore, evaluation of such behaviors should be carried out in single-compound systems, which are simpler to analyze than cement but feature the same effects. As noted previously, C<sub>3</sub>S is the dominant cement phase (comprising 50-70%<sub>mass</sub> of cement) [1]. The hydration of C<sub>3</sub>S produces two hydrates, that is, C-S-H and CH (portlandite), in stoichiometric quantities, and—like in cement pastes—the nucleation and growth

of C–S–H is the driving mechanism in  $C_3S$  pastes. As such,  $C_3S$  is deemed a simpler single-compound alternative for cement, and understanding PCE– $C_3S$  interaction can be the basis for understanding PCM–cement interactions.

In this study, a combination of experimental and simulation techniques is used to elucidate the effect of PCE on hydration of C<sub>3</sub>S. To fully examine and describe the links between PCE's molecular architecture and its ability to suppress hydration of C<sub>3</sub>S, PCEs with three different molecular architectures—albeit, of the same polymer family—are used. The hydration kinetics are monitored, using isothermal microcalorimetry technique, across a broad range of PCE dosages in C<sub>3</sub>S pastes. A modified pBNG simulation routine-which has only recently been applied in the literature [11, 25, 33]—is employed to reproduce, and subsequently describe, hydration kinetics of such systems. Focus is given to consolidate results obtained from experiments and simulations, and analyze them in tandem to elucidate the mechanistic origins of PCE-induced suppression of C<sub>3</sub>S hydration—including both early and later stages, wherein C<sub>3</sub>S hydration is driven by dissolution and nucleation-and-growth, respectively. The mechanisms are ultimately distilled into a single numerical equation that correlates the molecular architecture and dosage of PCE with its potential to influence C<sub>3</sub>S hydration kinetics. Such correlation is of significance for practical applications as it can be used as a robust, quantitative basis to compare and rank PCEs on the bases of their potential to suppress C<sub>3</sub>S hydration rates, and to select optimum ones based on the type/nature of the application.

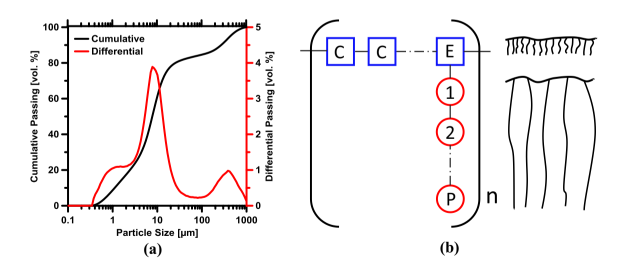
Table 1 Architectural parameters of the three PCEs—as determined from SEC-MALS, HPLC, and potentiometric titration techniques [20, 37]

PCE#	C/E	Р	п	M <sub>w</sub>
PCE-1	1.80	23	20.05	25,145.07
PCE-2	2.10	23	18.10	23,166.84
PCE-3	5.20	17	9.35	11,990.65

Here, *C/E* (unitless) is the ratio of the number of carboxylate functional groups to the number of ester functional groups, *P* (unitless) is the number of ethylene oxide monomers per side chain, *n* (unitless) is the number of side chains per PCE backbone (or the number of repeating units), and  $M_w$  (g mol<sup>-1</sup>) is the calculated molecular weight.  $M_w$  is equal to [*n*. {*P*.  $M_{w,SC} + (C/E + 1)$ .  $M_{w,BB}$ ]—wherein,  $M_{w,SC}$  (g mol<sup>-1</sup>) and  $M_{w,BB}$  (g mol<sup>-1</sup>) are the molecular weights of the side chain and backbone, respectively [20, 22]. The nomenclature of the architectural parameters of PCEs was adapted from previous studies [15, 20, 59]

#### 2 Materials and experimental methods

Triclinic C<sub>3</sub>S (T<sub>1</sub>–Ca<sub>3</sub>SiO<sub>5</sub>) was synthesized by following a high-temperature synthesis route using phase-pure precursor materials. Calcium oxide (CaO), the residual material left after calcite (CaCO<sub>3</sub>) decomposition, was mixed with silica (SiO<sub>2</sub>,  $\alpha$ -quartz) in stoichiometric proportion (CaO:SiO<sub>2</sub>=3:1), and pelletized at high pressure of 100 MPa. The pellets were then thermally treated in a furnace for 12 h at 1600 °C in platinum receptacles, and subsequently air-quenched [34, 35]. The sintered pellets were then crushed with mortar and pestle, and ground



**Fig. 1 a** The cumulative (primary y-axis) and differential (secondary y-axis) particle size distribution (PSD) of  $C_3S$ . The largest relative uncertainty in the median diameter ( $d_{50'}$  µm) of  $C_3S$ , based on six replicate measurements, was on the order of ±6%. **b** Schematic

representation of the comb-shaped architecture of PCE, wherein n is the number of repeating units, P is the side chain length, and C/E represents the carboxylate-to-ether ratio

in a ball mill for 24 h. The resulting powder was analyzed via X-ray diffraction (Panalytical X'Pert Pro MPD) and was determined to be C<sub>3</sub>S powder containing  $\approx 0.80\% \pm 0.20\%$  residual CaO by mass by Rietveld analysis. The particle size distribution (PSD: Fig. 1a) of C<sub>3</sub>S was measured using static light scattering analyzer (Microtrac S3500) [36]. The median particle size (d<sub>50</sub>, µm) of C<sub>3</sub>S particulates was determined as 7.78 µm. By combining the PSD of C<sub>3</sub>S with its density (i.e., 3150 kg m<sup>-3</sup>), the specific surface area (SSA<sub>C3S</sub>) of C<sub>3</sub>S particulates was calculated as 562 m<sup>2</sup> kg<sup>-1</sup>.

Three different polymeric, comb-shaped, PCEs (with solid mass contents of 30%) were used in this study (Fig. 1b). All three PCEs belong to the same polymer family—consisting of polymethacrylic backbone, grafted with polyethylene oxide side chains—albeit their molecular architectures are different. The different PCEs are subsequently referred to as PCE-1, PCE-2, and PCE-3. Details pertaining to the PCEs' molecular architecture, as determined from size exclusion multi-angle light scattering (SEC-MALS), high-performance liquid chromatography (HPLC) and potentiometric titration techniques [20, 37], are listed in Table 1.

Pastes were prepared by mixing deionized-water (DIwater) and C<sub>3</sub>S at a constant liquid-to-solid mass ratio (l/s) of 0.45. To describe the role of PCE on C<sub>3</sub>S hydration kinetics, the three different PCEs (i.e., PCE-1, PCE-2, and PCE-3, described in Table 1) were added to the pastes at dosages (C<sub>PCF</sub>) of 0.000, 0.625, 1.250, 1.875, and 2.500% (by mass of  $C_3S$ ). It is pointed out that these dosages signify the total (i.e., solid + liquid) mass of the PCE. Based on the liquid content of the admixture (i.e.,  $\approx 30\%_{mass}$  of all PCEs), the aforementioned dosages would amount to 0.000, 0.188, 0.375, 0.563, and 0.750% by solid component of the PCE per unit mass of the binder. The upper bound of dosage, that is, 2.500%, was determined by saturation point test [25, 38] for PCE-1 (with respect to cement paste), and is representative of dosages used in high-performance concretes [13]. The lower dosages of PCE correspond to 25, 50, and 75% of the upper bound, respectively. For provision of PCE into the paste, the mixing-protocol involved mixing of DI-water and PCE for 20 s, followed by an additional minute of mixing with C<sub>3</sub>S. For experiments where the upper bound of PCE dosage was employed, PCE was also deployed in delayed mode. Specifically, in delayed mode, a 5-min period, from when the DI-water first came into contact with  $C_3S$ , was allowed to elapse before PCE was introduced to the paste. Here, prior to the addition of PCE, the paste was mixed for 1 min, and for another 20 s after PCE was added.

 $C_3S$  hydration kinetics in pastes containing approximately 1 g of anhydrous  $C_3S$  was monitored for a minimum of 72 h (or 144 h for pastes containing PCE-3), at a constant temperature of 20 °C±0.01 °C, using a TAM IV isothermal microcalorimeter. Microcalorimetry techniques are able to monitor heat evolution, resulting from a chemical reaction, at a high resolution  $(10^{-8} \text{ J s}^{-1})$ . The differential and cumulative heat evolution profiles were divided (or normalized) by the enthalpy of C<sub>3</sub>S hydration [1, 35], that is, 484 J g<sub>C3S</sub><sup>-1</sup>, to determine the rate of hydration  $(da/dt, \text{ units of h}^{-1})$  and the degree of hydration (*a*, reaction mass fraction of C<sub>3</sub>S) of C<sub>3</sub>S, respectively, as functions of time. The values of *a* and *da/dt* calculated in such manner are premised on the assumption that the heat release, determined from microcalorimetry methods, is exclusively due to C<sub>3</sub>S hydration. In the context of experiments conducted in this study, the aforementioned assumption is reasonable because the heat release associated with physical and chemical interactions between PCE and C<sub>3</sub>S paste components is minuscule compared to heat released from the hydration of C<sub>3</sub>S [25, 39].

A thermogravimetric analyzer (TGA, SDT600) was used for identification and quantification of phases present in the binder after 24 h of hydration. Hydration was stopped by crushing the hydrated pastes, into small grains, immersing them in isopropanol for 24 h [40], followed by drying in the oven (T=85 °C) for an additional 24 h. The samples were then ground into fine powder. The powder samples were heated in an inert atmosphere of N<sub>2</sub> over a temperature range of 30–900 °C. The cumulative and differential mass loss traces were used to quantify the amount of CH present in the system; towards this, well-established methods detailed in prior studies [40, 41] were used.

#### 3 Phase boundary nucleation and growth model

A modified pBNG model is applied to describe the influence of PCE on early-age hydration kinetics of C<sub>3</sub>S. Akin to classical pBNG models applied to cementitious systems [3, 7, 8, 42–45], the model used herein assumes that a single product of constant density forms heterogeneously on solidphase substrate boundaries (i.e., C<sub>3</sub>S particles' surfaces) at a given nucleation event (i.e., at time =  $\tau$  h), and assumes that its subsequent growth drives and controls—as the rate controlling mechanism—the kinetics of C<sub>3</sub>S hydration. This assumption—typically termed as *site saturation*—entails that after the initial burst of nucleation, no further nuclei of the product form. As per these criteria, the volume fraction of the product within the paste [*X*(*t*), unitless] is given by Eq. 11 [1, 25, 33, 35, 42, 46].

$$X(t) = 1 - exp\left[-2k_G \cdot (t-\tau) \cdot \left(1 - \frac{F_D[k_s \cdot (t-\tau)]}{k_s \cdot (t-\tau)}\right)\right]$$
(1)

where, 
$$F_D(x) = exp(-x^2) \int_0^x exp(y^2) dy$$
 (2)

$$k_{s} = G_{out}(t) \cdot \left(\pi \cdot g \cdot I_{density}\right)^{\frac{1}{2}}$$
(3)

$$g = \left[\frac{G_{par}(t)}{G_{out}(t)}\right]^2 \tag{4}$$

$$k_G = r_G \cdot G_{out}(t).a_{BV} \tag{5}$$

In Eq. 1,  $F_D$  represents the f-Dawson function shown as the integral in Eq. 2. The parameter  $k_s$  (h<sup>-1</sup>) represents the inverse of time needed by the product to completely cover the surface of the anhydrous  $C_3S$  particles [7, 42] (Eq. 3). Its value depends on the nucleation density of the product  $(I_{density} \mu m^{-2})$ , that is, the number of product nuclei per unit surface area of C<sub>3</sub>S particulates, as well as the product's growth rate and geometry. In this study, it is assumed that the growth of the product occurs in an anisotropic manner, while varying with respect to time. The growth rates in the outward (i.e., normal to and away from C<sub>3</sub>S particles' surface) and lateral (i.e., parallel to the C<sub>3</sub>S particles' surface boundary) directions are represented as G<sub>out</sub> (t) and  $G_{par}(t)$ , respectively. Along the two-dimensional plane parallel to the C<sub>3</sub>S surface,  $G_{par}(t)$  is assumed to be isotropic [7, 25, 33]. It is worth noting that such temporal variation of the product growth rate is a departure from classical pBNG models—wherein, throughout the entire duration of hydration, the growth rate is assumed to remain constant. This implementation of variable product growth rate—based on the original study of Bullard et al. [46], and subsequently adopted by several researchers [11, 25, 33, 35]—captures sharp changes in C–S–H's growth rate as its supersaturation in the solution varies in a highly nonlinear fashion with time. While G<sub>out</sub> and G<sub>par</sub> vary with time, a ratio of 1:0.50 for G<sub>out</sub>:G<sub>par</sub> is maintained; as such, the anisotropy factor [i.e., g (unitless), shown in Eq. 4] of the product nuclei, remains constant at 0.25 throughout the entirety of C<sub>3</sub>S hydration measured via microcalorimetry. This relationship between  $G_{out}$  and  $G_{par}$  causes the product to acquire aspherical geometry [7, 42]—essentially mimicking fiber-like geometry of C-S-H observed experimentally at early ages [47, 48].

In Eq. 1, the parameter  $k_G$  (h<sup>-1</sup>) represents the inverse of time required for occupation of the capillary pore volume by the product.  $k_G$  is a function of  $G_{out}$ , and another constant  $r_G$  (unitless); here,  $r_G$  (Eq. 5) represents the ratio of product growth rate into vis-à-vis out of the substrate in the direction normal to the substrate. In previous studies [7, 42], focused on early age hydration of cement (and  $C_3S$ ), it has been reported that hydrates do not penetrate the anhydrous particles, and, therefore,  $r_G \approx 0.50$ . As simulations presented in the current study pertain to early ages, the value of  $r_G$  is assumed to be constant at 0.50. The value of  $k_{G'}$  and thus kinetics of  $C_3S$  hydration, also depend on the boundary area of the substrate per unit volume of the paste  $(a_{BV}, \mu m^{-1})$  (Eq. 6). The paste's volume is simply the initial cumulative volume of the paste (i.e., volumes of C<sub>3</sub>S and water), and the substrate's area is the initial surface area of C<sub>3</sub>S particles that is available for nucleation of product. In Eq. 6,  $SSA_{C3S}$  (units of m<sup>2</sup> kg<sup>-1</sup>) is the specific surface area of C<sub>3</sub>S particles,  $\rho_W$  is the density of water (i.e., 1000 kg m<sup>-3</sup>) and  $\rho_{C3S}$  is the density of C<sub>3</sub>S (i.e., 3150 kg m<sup>-3</sup>). By combining  $SSA_{C3S}$  and  $I_{density}$  the total number of supercritical product nuclei ( $N_{nuc}$ , unit of  $g_{C3S}^{-1}$ ) produced per gram of C<sub>3</sub>S can be calculated [33, 43, 45, 49] (Eq. 7).

$$a_{BV} = \left[\frac{SSA_{C_3S}}{\left\{\frac{\frac{l}{s}}{\rho_W} + \frac{1}{\rho_{C_3S}}\right\}}\right]$$
(6)

$$N_{nuc} = SSA_{C3S} \cdot I_{density} \tag{7}$$

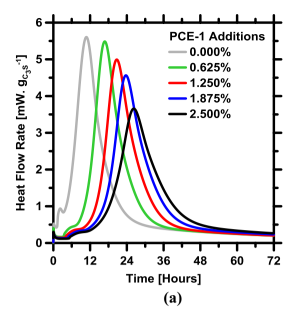
The fraction of the paste' volume occupied by the product [X(t)], as calculated from Eq. 1, and the degree of hydration (*a*) of C<sub>3</sub>S are linked by yet another constant, *B* (unitless) [25, 35, 46], as shown in Eqs. 8–9:

$$\alpha(t) = B \cdot X(t) \tag{8}$$

$$B = \left[ \left( \frac{\frac{\rho_{C_3S}}{\rho_{products}}}{\frac{\frac{l}{s} \cdot \rho_{C_3S}}{\rho_{water}} + 1} \right) \left( \frac{C + \frac{1}{\rho_{C_3S}} - \frac{1}{\rho_{water}}}{\frac{1}{\rho_{product}} - \frac{1}{\rho_{water}}} \right) \right]^{-1}$$
(9)

where,  $\rho_{product}$  is the average wet density of hydrates (assumed to be 2070 kg m<sup>-3</sup>, based on the stoichiometric ratio of formation and individual densities of the two hydrates, C–S–H and CH) [50, 51], and the parameter  $c=-7.04 \times 10^{-5}$  m<sup>3</sup> kg<sup>-1</sup> stands for the chemical shrinkage of the paste that occurs when 1 kg of C<sub>3</sub>S is fully hydrated [35, 46, 52].

Based on the above equations (i.e., Eqs. 1–9), to numerically reproduce the experimentally-derived reaction rates, the variables that need to be ascertained are:  $G_{out}$ (t) and  $I_{density}$ . Of these two variables,  $G_{out}(t)$  is a function of time, whereas  $I_{density}$  is constant (with respect to time). For a given system, to determine the optimum functional form of  $G_{out}$  and the optimum value of  $I_{density}$  and, a Nelder–Mead based simplex algorithm [35, 53, 54], based on non-linear optimization and derivative-free routines, is implemented in two steps. In the first step, the value of  $G_{out}$  is kept constant at 0.075 µm h<sup>-1</sup>—a value derived from microscopy-based analyses of early age C–S–H growth in C<sub>3</sub>S and similar systems [47, 48, 55]. The algorithm varies the values of  $I_{density}$  within predefined bounds (i.e.,



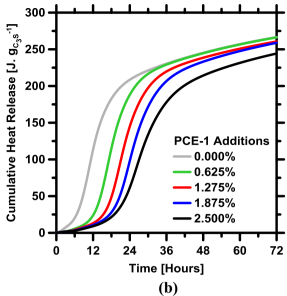


Fig. 2 Isothermal microcalorimetry based determinations of timedependent **a** heat flow rate and **b** cumulative heat release of  $C_3S$  pastes provisioned with PCE-1 at different dosages ( $C_{PCF}$ ). The I/s for

0.01-to-100  $\mu$ m h<sup>-1</sup>) until the deviation between measured and simulated rates of reaction (da/dt) is minimalized. It is worth highlighting that up to the first simulation step, the model represents the conventional pBNG formulation [8]—wherein the anisotropic growth of product, several nuclei of which precipitate at a virtual time  $\tau$  (h), is assumed to be constant. To factor in the temporal variation in product growth rate, the second and final simulation step is employed. In this step, at a given time t, the optimum value of I<sub>density</sub>, determined from the first step, is used as constant, whereas  $G_{out}$  is iteratively varied between  $10^{-3}$ and  $10^3 \,\mu\text{m} \,\text{h}^{-1}$ . At convergence, that is, when the deviation between the simulated and measured reaction rates reaches its minimum (i.e., within 0.05%), the value of G<sub>out</sub> yielded by the optimization process is finalized as the optimum. By implementing the optimization process over 72 h (or longer for PCE-3 containing pastes) of hydration, using a time step of 0.01 h, the optimum values of  $G_{out}$  for the entire duration of C<sub>3</sub>S hydration are thus determined. The functional form of G<sub>out</sub>, obtained from the optimization routine, mimics the product's non-monotonic and nonlinear evolution of growth rate as a function of its supersaturation in the solution. In prior publications [11, 25, 33, 35, 46], which employ similar simulation scheme, it has been shown that such functional form of the product growth rate—as obtained from the simulations—reproduced the intrinsic changes in the evolution of the solution's chemistry (e.g., changes in pH, ionic strength, and water activity). Therefore, whereas this scheme of deriving the functional form of the product growth rate is indirect, the final results

pastes included in these figures, as well as the subsequent ones, is 0.45. The uncertainty in measured heat flow rate at the main hydration peak is  $\pm\,2\%$ 

are still reflective of the physical processes occurring in the system [11, 35, 46, 48, 55].

#### 4 Results and discussion

Figure 2 shows representative heat evolution profiles of  $C_3S$  pastes provisioned with PCE-1 at different dosages (i.e., 0–2.5%).

As can be seen, PCE significantly suppresses C<sub>3</sub>S hydration rates, as marked by various characteristic aspects of the pastes' heat evolution profiles: lengthening of the induction period (the period between the initial wetting peak and the onset of acceleration), rightward shift of the heat evolution curves, and reduced heat flow rates at the main hydration peak. The suppression of hydration induced by PCE increases monotonically with its dosage, which entails good correlation between amount of PCE present in the paste and the resultant reduction of C<sub>3</sub>S hydration rates. This correlation is indicative of inhibition of C<sub>3</sub>S dissolution sites (thus causing prolongation of the induction period), and C-S-H nucleation sites (thus causing slower approach to the main hydration peak)—both of which most likely manifest as a result of adsorption of PCE molecules on C<sub>3</sub>S particles' surfaces, and scale with PCE dosage [19, 22–25]. It is worth pointing out that provision of PCE in the paste retards not only the approach to the main hydration peak but also the departure from it (i.e., slightly lower slope of the deceleration regime, as shown in Fig. 2a). Slower post-peak deceleration in  $[C_3S + PCE]$ 

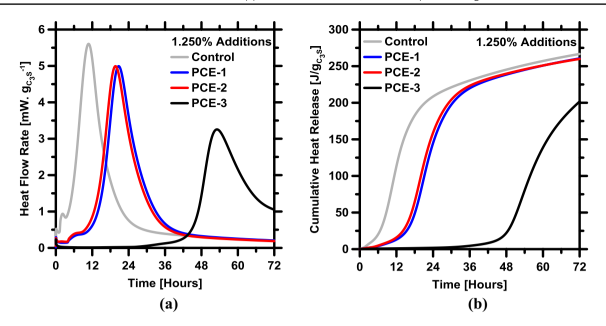


Fig. 3 Isothermal microcalorimetry based determinations of time-dependent **a** heat flow rate and **b** cumulative heat release of  $C_3S$  pastes provisioned with different PCEs (i.e., PCE-1, PCE-2, and PCE-3) at equivalent dosage ( $C_{PCF}$ ) of 1.25%

pastes implies that, at later ages, the rate of hydration of C<sub>3</sub>S in such pastes is relatively faster than in the control system (i.e., [C<sub>3</sub>S+0% PCE] paste); as such, much of the loss in early-age reactivity, that is induced by PCE, is recouped at later ages. This is better shown in Fig. 2b, wherein, at later ages (i.e.,  $\approx$ 72 h), the cumulative heat release of [C<sub>3</sub>S+PCE] pastes converge—or, appear to be on track to converge—with that of the control system.

Results shown in Fig. 2, and the above discussion, describe the role of PCE dosage on  $C_3S$  hydration rates. To better understand the effects of PCE's molecular architecture on  $C_3S$  hydration kinetics, heat evolution profiles of  $[C_3S + PCE]$  pastes, prepared with equivalent dosages but different types of PCE, were compared (Fig. 3).

It is evident that regardless of the molecular architecture, all three PCEs suppress  $C_3S$  hydration at early ages (Fig. 3a). In addition to the pre-peak suppression of hydration (e.g., slower approach to the main hydration peak), the slower deceleration beyond the main hydration peak is common among the three PCEs. The slower deceleration results in convergence (or, in case of PCE-3, a trajectory that would eventually result in convergence) of cumulative heat release at later ages. While, qualitatively, the general nature of hydration suppression is similar amongst the three PCEs, there are substantial differences in the magnitude of such decelerations. As can be seen in Fig. 3,  $C_3S$  hydration is more significantly suppressed by PCE-3 as compared to PCE-1 and PCE-2; the

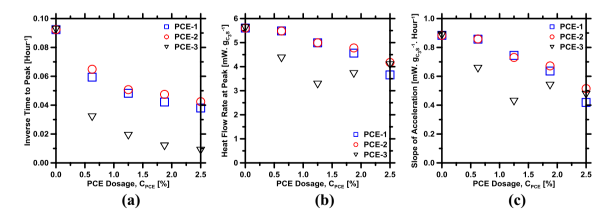


Fig. 4 The calorimetric parameters: **a** inverse of time to the main hydration peak, **b** heat flow rate at the main hydration peak, and **c** slope of the acceleration regime, extracted from heat evolution

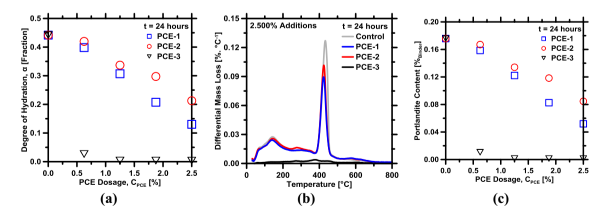
profiles, and plotted against PCE dosage. For a given system, the uncertainty in each calorimetric parameter is  $\pm\,2\%$ 

latter two produce similar magnitudes of suppression. Although the results shown in Fig. 3 pertain to a single PCE dosage (i.e., 1.25%), the stark difference—between the magnitude of suppression of C<sub>3</sub>S hydration induced by PCE-3 vis-à-vis those by PCE-1 and PCE-2-was also observed at other dosages. Since all PCEs belong to the same polymer family (i.e., same composition of backbone and side chains), it is clear that the differences (or similarities) in their potential to suppress C<sub>3</sub>S hydration arise due to intrinsic differences (or similarities) in their molecular architecture. Such architectural differences are expected to dictate their adsorption capacity, and, thus, their ability to inhibit sites of C<sub>3</sub>S dissolution and C-S-H nucleation. As PCE-3 has the highest potential to suppress C<sub>3</sub>S hydration, it can be said that its molecular architecture is more favorable towards adsorption on C<sub>3</sub>S particles' surfaces. Along the same lines, as PCE-1 and PCE-2 produce equivalent suppressions of hydration at equivalent dosages, it is expected that their molecular architectures, and thus their adsorption potentials, are broadly similar. Further details pertaining to the role of PCE's molecular architecture on C<sub>3</sub>S hydration suppression are discussed later in this section.

To better contrast suppression of  $C_3S$  hydration, as prompted by the three PCEs, characteristic calorimetric parameters [i.e., inverse of time corresponding to the main hydration peak (h<sup>-1</sup>), heat flow rate at the main hydration peak (mW  $g_{C3S}^{-1}$ ), and slope of the acceleration regime (mW  $g_{C3S}^{-1}$  h<sup>-1</sup>]—indicative of acceleration or retardation in hydration kinetics [25, 33, 43, 56, 57]—were extracted and plotted against the PCE dosage (Fig. 4).

As can be seen, with increasing PCE dosage, the induction period's length increases (Fig. 2); this delays the incidence of the main hydration peak (Fig. 4a). This indicates that interactions between PCE and  $C_3S$  delay the hydrate nucleation event (i.e., massive precipitation of C–S–H and CH, which occurs around the time when the induction period terminates) [2, 9, 25, 58]. These results are in good agreement with prior studies [17, 22, 24, 25], which have reported that the adsorption of PCE molecules on C<sub>3</sub>S particles' surfaces blocks C<sub>3</sub>S dissolution sites, which causes deceleration of C<sub>3</sub>S dissolution and—as a consequenceprolongs the induction period. Akin to the trends in (inverse of) time of the main hydration peak, the other two calorimetric parameters—heat flow rate at the peak and slope of the acceleration regime-also decrease progressively with increasing amount of PCE in the paste (Fig. 4b, c). These results suggest that PCE not only delays the time of product nucleation but also suppresses the product's post-nucleation precipitation rate—as can also be seen in Figs. 2 and 3. Past literature [24, 25] suggests that such delay occurs due to interactions between PCE and C-S-H nuclei. More specifically, these studies [24, 25] suggest that PCE molecules (i.e., those that are not adsorbed on C<sub>3</sub>S particles' surfaces and still remain in the solution) adsorb onto positively charged C-S-H surfaces, partially blocking their access to the contiguous solution, and, thus, diminishing their growth rate (and, hence, their precipitation rate). Furthermore, as PCE molecules remain adsorbed on C-S-H surfaces, the aforementioned inhibition of C-S-H's growth persists even at later stages of hydration. This is better revealed in Figs. 2a and 3a, wherein it is shown that the post-peak deceleration of [C<sub>3</sub>S + PCE] paste is slower than in the control paste. Due to this, the cumulative heat release of  $[C_3S + PCE]$  pastes approach convergence with that of the control system at later ages (Figs. 2b, 3b).

The equivalency in calorimetric parameters of pastes prepared with PCE-1 and PCE-2 (Fig. 4) suggests that the hydration-suppression effects of the two PCEs are broadly similar. This equivalency is corroborated in Fig. 5, which shows that in  $[C_3S+PCE-1]$  and  $[C_3S+PCE-2]$  pastes—when prepared with equivalent dosage of PCE—the degree of



**Fig. 5 a** Isothermal microcalorimetry based determinations of time-dependent degree of reaction ( $\alpha$ ) of C<sub>3</sub>S. **b** DTG traces showing differential mass loss profiles, and **c** mass contents of portlan-

dite (CH: as  $\%_{mass}$  of the binder) as determined from analyses of DTG traces, in [C<sub>3</sub>S+PCE] pastes at 24 h. The highest uncertainty in phase quantifications by DTG is  $\pm\,2.5\%$ 

hydration of  $C_3S(\alpha)$  and portlandite (CH) contents at 24 h are similar. These results are not surprising because PCE-1 and PCE-2 have similar molecular architectures-with similar values of C/E, n, P, and  $M_{w}$  (see Table 1)—and, thus, are expected to have similar potentials to adsorb on silicate (i.e., C<sub>3</sub>S and C-S-H) surfaces and similar capacities to block their access to the contiguous solution. Comparisons between PCE-1/PCE-2 and PCE-3 are, however, not trivial because of significant differences in their molecular architectures (Table 1). As can be seen in Fig. 5, at a given dosage, suppression of hydration induced by PCE-3 is considerably more severe (i.e., lower  $\alpha$  and lower CH content at 24 h) as compared to those induced by PCE-1 and PCE-2. As the C/E of PCE-3 is significantly larger (i.e., C/E = 5.20) as compared to those of PCE-1 and PCE-2 (i.e., C/E = 1.80 and 2.10, respectively), it can be said that the lower side chain grafting density of PCE-3 permits a higher negative charge on the backbone, and, thus, improved adsorption onto positively charged silicate surfaces [15, 26–28]; this manifests as greater suppression of hydration in [C<sub>3</sub>S + PCE-3] pastes. Furthermore, the length of the side chain (i.e., given by P) of PCE-3 is shorter (i.e., P = 17) than those of PCE-1 and PCE-2 (i.e., P = 23). As prior studies [17, 29] have indicated, smaller side chains ensure that access to the negative charges on the PCE's backbone is not hindered or limited by steric hindrance (induced by the side chains). This enables better adsorption of PCE molecules onto silicate surfaces, thus resulting in superior hydration inhibition. Lastly, on account of lower side chain grafting density, shorter side chain length, and smaller number of repeating units (i.e., n), the overall molecular weight of PCE-3 is lower (i.e.,  $M_w = 11,990$  g mol<sup>-1</sup>) than PCE-1 and PCE-2 (i.e.,  $M_w = 25,145$  and 23,166 g mol<sup>-1</sup>). This entails that for a given (mass-based) dosage, the number of PCE-3 molecules in the paste is higher than those prepared with PCE-1 or PCE-2. On account of their larger population in the paste, PCE-3 molecules are able to adsorb more effectively on C<sub>3</sub>S and C–S–H surfaces, and, thus, suppress C<sub>3</sub>S hydration to a greater extent as compared to the other two PCEs.

The discussion in the above paragraph provides a basis—albeit qualitative—for explaining the links between molecular architecture of PCE and its ability to suppress C<sub>3</sub>S hydration. However, through such qualitative basis, it is impossible to quantify the superiority of a given PCE—in terms of inducing retardation of C<sub>3</sub>S hydration kinetics—as compared to another PCE of a different molecular architecture. For example, while it is known that higher and lower values of C/E and P, respectively, cause superior suppression of C<sub>3</sub>S hydration, such knowledge does not allow quantitative prediction of how much more retardation a PCE (say, with higher C/E and lower P compared to another PCE) will induce compared to another. In a recent paper, Marchon et al. [59] suggested that the net retardation of C<sub>3</sub>S (or cement) hydration kinetics induced by PCE is proportional to: (1) the concentration of PCE molecules—expressed as the ratio of its dosage in paste ( $C_{PCE}$ : expressed as  $\%_{mass}$ ) to its molecular weight  $(M_w)$ —and (2) a function (f) of C/E, shown in Eq. 10. The former proportionality (i.e., with respect to PCE concentration) is expected because larger number of PCE molecules in the paste would entail greater probability of their adsorption on silicate (i.e., C<sub>3</sub>S and C-S-H) surfaces, and, therefore, greater retardation. It is worth noting that PCE concentration inherently incorporates the effects of dosage and architectural parameters of PCE (i.e., n, P, C/E, and molecular weights of the side chain and backbone—as shown in Table 1 and described in [20, 22]), and, therefore, serves as an encompassing parameter on its own. The latter proportionality [i.e., with respect to f(C/E)] captures

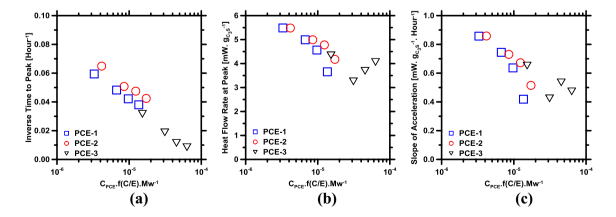


Fig. 6 The calorimetric parameters: **a** inverse of time to the main hydration peak, **b** heat flow rate at the main hydration peak, and **c** slope of the acceleration regime extracted from the calorimetry

profiles, and plotted against the composite architectural parameter of the PCE ( $P_{PCE}$ : calculated using Eq. 11). The uncertainty in the value of  $P_{PCE}$  is on the order of ± 15%

the dependency of adsorption potential of PCE on the strength of electric field induced by its molecules on the adsorbent's surface. As described in prior publications [20, 59, 60], higher value of f(C/E) implies greater magnitude, better exposure, and wider distribution of electric field induced by the PCE, and, therefore, better adsorption of its negatively charged backbone on the positively charged silicate surfaces. Marchon et al. [59] proposed that the overall retardation of C<sub>3</sub>S (or cement) hydration kinetics, as induced by a given PCE, could be quantified by consolidating the aforementioned proportionalities into a composite architectural parameter ( $P_{PCF}$ : expressed in arbitrary units), as shown in Eq. 11. It must be noted that Marchon et al. [59] originally defined  $P_{PCE}$  using the molar mass of the repeating unit in the denominator. However, in Eq. 11, the molecular weight of  $(M_w)$  the entire PCE molecule is used (instead of just the repeating unit) in order to account for the PCE's backbone's length in addition to other architectural features. In Eq. 11, the variables (i.e., C<sub>PCF</sub>, C/E and  $M_{w}$ ) are expressed in their typical units as described in the **Experimental Section.** 

$$f(C/E) = \left(\frac{C/E}{C/E+1}\right)^{\frac{3}{2}}$$
(10)

Composite Architectural Parameter ( $P_{PCE}$ )

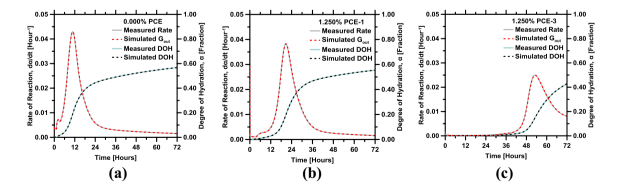
$$=\frac{c_{PCE}}{M_w}f(C/E)=\frac{c_{PCE}}{M_w}\left(\frac{C/E}{C/E+1}\right)^{\frac{1}{2}}$$

To test the efficacy of Eq. 11, in terms of predicting the retardation caused by PCE, the calorimetric parameters—which capture alterations in C<sub>3</sub>S hydration kinetics (as shown in Fig. 4)—were plotted against the composite architectural parameter ( $P_{PCE}$ ) of PCE. As can be seen in Fig. 6, in spite of substantial differences in dosages and molecular architectures of PCEs, each calorimetric parameter—extracted from heat evolution profiles of different pastes—converges, broadly, into a unified master trend.

Admittedly, the datapoints of two of the calorimetric parameters, that is, peak heat flow rate (Fig. 6b) and slope of the acceleration regime (Fig. 6c), are not as convergent as in the case of inverse of time to peak (Fig. 6a). These minor deviations—from the unified master trend—can be attributed to errors associated with the experimentallydetermined parameters (e.g., PCE architectural parameters such as C/E, and calorimetric parameters). In spite of the aforementioned deviations, in general, it can be said that the deceleration of C<sub>3</sub>S hydration kinetics increases, broadly in a logarithmic manner, with respect to increasing values of the composite architectural parameter ( $P_{PCF}$ ) of PCE. This correlation suggests that Eq. 11 can be used as a robust, quantitative basis to compare and rank PCEs on the basis of their potential to suppress C<sub>3</sub>S hydration rates. Equation 11 also allows prediction of additional (or reduction in) retardation-with respect to a benchmark system—if the PCE's dosage or molecular architecture are altered. As an example, if retardation of C<sub>3</sub>S hydration caused by a given PCE is measured (e.g., using calorimetry), additional retardation at a higher dosage of the same PCE (or, a different PCE) can be readily predicted by plugging in the PCE dosage (or, architectural parameters) in Eq. 11.

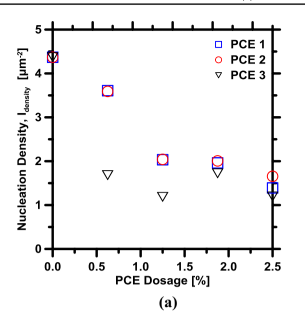
The experimental results discussed thus far shed light on the effects of PCE dosage and molecular architecture on the overall hydration kinetics of  $C_3S$ . To better understand such effects, specifically in the context of alterations in nucleation and growth of the main hydrate (i.e., C–S–H), the pBNG model was applied. Figure 7 shows the simulated reaction rates of pastes compared against those obtained from experiments.

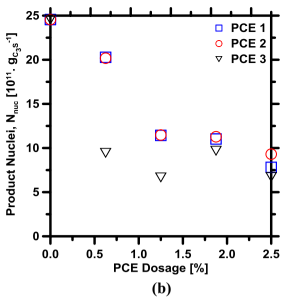
As can be seen, through optimization of the outward growth rate of the product  $[G_{out}(t)]$  and the product



**Fig. 7** Representative set of simulated and measured reaction rates  $(d\alpha/dt; primary y-axis)$  and degree of reaction ( $\alpha$ ; secondary y-axis) for C<sub>3</sub>S pastes prepared with varying PCE architectures and dos-

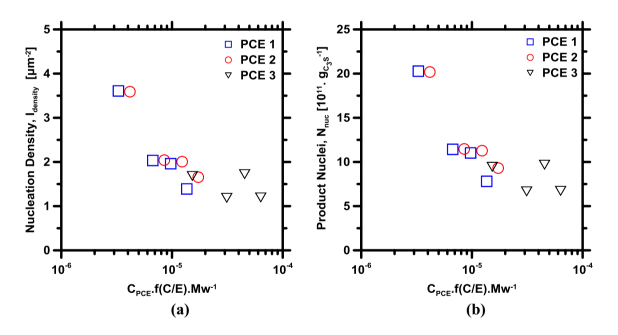
ages of: **a** 0.000% PCE, **b** 1.250% PCE-1, and **c** 1.250% PCE-3. Simulations are deterministic, and, therefore, there is no uncertainty associated with them





**Fig. 8** Parameters derived from pBNG simulations: **a** nucleation density of the product ( $I_{density}$ ), and **b** total number of super product nuclei formed per gram of C<sub>3</sub>S ( $N_{nuc}$ : calculated from Eq. 7), plotted

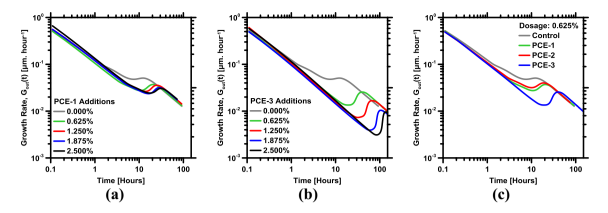
against PCE dosage ( $C_{PCE}$ ). Simulations are deterministic, and, therefore, there is no uncertainty associated with them



**Fig. 9** Parameters derived from pBNG simulations: **a** nucleation density of the product ( $I_{density}$ ), and **b** total number of super product nuclei formed per gram of C<sub>3</sub>S ( $N_{nuc}$ ), plotted against the composite architectural parameter of PCE ( $P_{PCE}$ : calculated from Eq. 11)

nucleation density (*I*<sub>density</sub>), the experimental results are well reproduced by the model. Variations in these simulation parameters are investigated below to delineate the mechanistic origins of modifications in the nucleation and growth process in relation to PCE's molecular architecture and dosage.

Based on optimization of parameters by the pBNG model, it was found that PCE has profound effect on the hydrate nucleation event, which occurs at early ages, that is, in proximity to the time of termination of the induction period. As can be seen in Fig. 8a, the product nucleation density decreases monotonically with increasing PCE dosage. This manifests as a progressive



**Fig. 10** The temporal evolution of outward growth rate of the product ( $G_{out}$ ), as obtained from pBNG simulations, for: **a** pastes prepared with PCE-1 at different dosages, **b** pastes prepared with PCE-3 at different dosages, and **c** pastes prepared with different different with different dosages.

decline in the number of supercritical product nuclei  $(N_{nuc}: \text{calculated from Eq. 7})$ —forming heterogeneously on C<sub>3</sub>S surfaces—with respect to increasing PCE dosage (Fig. 8b). These results corroborate the hypotheses presented above, as well as those advanced in previous studies [24, 25]—that adsorption of PCE molecules on C<sub>3</sub>S surfaces inhibits topographical sites of C–S–H nucleation. As would be expected, at higher PCE dosages, larger fraction of C<sub>3</sub>S particles' surface is blocked, and, thus, a greater number of C–S–H nucleation sites are inhibited. It is interesting to note that at any given dosage, PCE-3 results in lower product nucleation density (Fig. 8a)—and, therefore, smaller number of product nuclei (Fig. 8b)—as compared to the other two PCEs.

ent PCEs at a fixed dosage of 0.625%. Simulations parameters are deterministic, and, therefore, there is no uncertainty associated with them

These results are in very good agreement with the trends derived directly from the experiments (Fig. 4), and suggest that the adsorption capacity—and, therefore, the potential to suppress C–S–H nucleation (and C<sub>3</sub>S hydration)—of PCE-3 is superior. As stated previously, the superior retardation induced by PCE-3 can be attributed to its intrinsic molecular architecture [namely, lower side chain grafting density (i.e., higher *C/E*), smaller side chain length (i.e., lower *P*), and lower molecular weight (i.e.,  $M_w$ )]—which enables better adsorption of its molecules onto C<sub>3</sub>S surfaces, and, therefore, more effective blocking of topographical C–S–H nucleation sites.

Earlier in this section, it was shown that the composite architectural parameter of PCE ( $P_{PCE}$ : calculated using

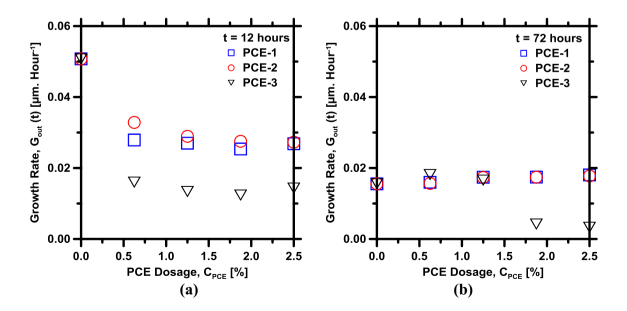


Fig. 11 The outward growth rate of the product, as obtained from pBNG simulations, at  $\mathbf{a} = 12$  h, and  $\mathbf{b} = 72$  h

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Eq. 11) correlated very well with the resultant effect on  $C_3S$  hydration (Fig. 5). To further verify the applicability of the composite parameter—in terms of capturing the combined effects of PCE dosage and molecular architecture—the product nucleation density and the number of product nuclei were plotted against  $P_{PCE}$ . As can be seen in Fig. 9, akin to the calorimetric parameters (Fig. 5), each pBNG parameter (i.e.,  $I_{density}$  and  $N_{nuc}$ ) converges onto a unified master trend; wherein, with increasing dosages of  $P_{PCE}$ , both  $I_{density}$  and  $N_{nuc}$  decrease monotonically.

It is clarified that the master trends, which emerge in Fig. 9, are not devoid of deviations. These deviations are, however, minor, and can be attributed to the statistical variance in the experimentally-derived parameters (e.g., C/E of PCE). Notwithstanding, these results—in conjunction with those shown in Fig. 5—provide compelling evidence that the composite architectural parameter of PCE ( $P_{PCE}$ ) is a reliable, and more importantly, a readily quantifiable indicator of the PCE's potential to suppress C<sub>3</sub>S hydration.

Finally, the product's outward growth rate  $[G_{out}(t)]$ , as obtained from pBNG simulations, are shown in Fig. 10.

As can be seen,  $G_{out}$  decreases by about three orders of magnitude—while evolving in a highly nonlinear manner with respect to time—over the course of C<sub>3</sub>S hydration. Such temporal evolution of the growth rate has been reported [25, 33, 35, 46] to mimic the time-dependent evolution of C–S–H supersaturation in the solution. It is noted that at any given time,  $G_{out}$ —particularly between ages of 1 h and 24 h—is lower at higher PCE dosages (Fig. 10a). Also, at equivalent dosages, Gout is lower in pastes prepared with PCE-3 as compared to those prepared with PCE-1 and PCE-2. To better contrast the influence of the different PCEs, values of G<sub>out</sub> were extracted at different times, i.e., at 12 h (early age), and at t = 72 h (later age), and, ultimately, plotted against the PCE dosage. As can be seen in Fig. 11a, at early ages,  $G_{out}$  in [C<sub>3</sub>S + PCE] pastes is consistently lower than in control pastes. This corroborates the hypothesis made earlier in this section-that the excess PCE molecules in the solution (that do not adsorb onto C<sub>3</sub>S particles' surfaces) adsorb onto C-S-H nuclei, and, subsequently, block their access to the adjacent solution, thus inhibiting their growth rate [24, 25]. At any given PCE dosage, lower values of  $G_{out}$  in  $[C_3S + PCE-$ 3] pastes, as compared to those prepared with other two PCEs, can be attributed to higher adsorption capacity of PCE-3 (i.e., on account of its molecular architecture), as has been explained previously in this section and highlighted in Figs. 4 and 8. At later ages, that is, at t = 72 h (Fig. 11b), the variations in Gout with respect to PCE's dosage and molecular architecture are significantly less pronounced. This is attributed to depletion of C–S–H supersaturation in the solution—which, in turn, causes G<sub>out</sub> to diminish [11, 24, 25, 33] and converge to similar values, regardless of dosage or molecular architecture of the PCE in the paste.

By and large, the results described so far support the theory that hydration of  $C_3S$  is suppressed by PCE—wherein, adsorption of PCE molecules onto  $C_3S$  surfaces suppresses  $C_3S$  dissolution and C–S–H nucleation. The adsorption of PCE molecules onto C–S–H surfaces

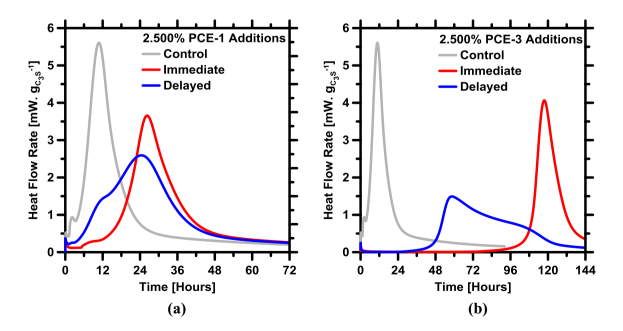


Fig. 12 Measured heat flow rates of  $[C_3S + 2.5\%$  PCE] pastes, prepared with **a** PCE-1, and **b** PCE-3. Results corresponding to both immediate and delayed (i.e., by 5-min) addition of PCE are shown. For comparison, the control system with no PCE is also shown

suppresses the post-nucleation growth of C-S-H. Based on this theory, it is hypothesized that if the adsorption of PCE onto C<sub>3</sub>S surfaces is reduced, the overall impact on inhibition of C<sub>3</sub>S dissolution and C-S-H nucleation sites would consequently be reduced. To test this hypothesis, additional experiments were conducted, wherein PCE was incorporated into the paste in delayed mode (i.e., addition of PCE five mins after mixing of C<sub>3</sub>S and DI-water). As can be seen in Fig. 12, when the addition of PCE is delayed, its impact on dissolution and nucleation site inhibition are indeed marginalized (albeit, not nullified). This manifests as shorter induction period and faster incidence of the main hydration peak compared to those in pastes provisioned with PCE at the time of mixing. Notwithstanding, in the case of delayed addition, the adsorption of PCE molecules onto surfaces of C-S-H-that form prior to, as well as after, the addition of PCE-results in substantial inhibition of its post-nucleation growth. This manifests as slower approach to and departure from the main hydration peak. It is also worth noting that in the case of delayed addition, as PCE molecules predominantly adsorb onto C-S-H surfaces (as opposed to  $C_3S$  and C-S-H surfaces), the suppression of post-nucleation growth of C-S-H (and, therefore, C<sub>3</sub>S hydration) is more profound compared to pastes in which PCE is added at the time of mixing (Fig. 12). The results shown in Fig. 12, and the mechanisms described above, are in good agreement with prior studies [24, 59].

# **5** Conclusions

A hierarchical sequence of experiments and pBNG simulations were employed to elucidate the effects of combshaped polycarboxylate ether (PCE) polymer on hydration mechanisms of tricalcium silicate ( $C_3S$ ). Emphasis was given to describe contributions of dosage and molecular architecture of PCE on early hydration of  $C_3S$ .

Results clearly show that hydration of  $C_3S$  is suppressed in presence of PCE—wherein, the deceleration scales with PCE content in the paste. The origin of such deceleration was hypothesized to be linked to the adsorption of PCE molecules on  $C_3S$  particles' surfaces, which inhibits topographical dissolution and C–S–H nucleation sites, and results in prolongation of the induction period. Furthermore, results suggest that adsorption of PCE molecules onto surfaces of C–S–H results in suppression of its postnucleation growth long after termination of the induction period. This results in a slower approach to, as well as departure from, the main hydration peak.

Through rigorous analyses of decelerating effects induced by three different PCEs, this study develops a robust correlation between the molecular architecture of PCE and its potential to suppress  $C_3S$  hydration. Results show that PCEs with lower side chain grafting density (i.e., higher *C/E*: carboxylate-to-ether ratio), smaller side chain length (i.e., smaller *P*: number of monomers per side chain), and lower overall molecular weight have greater potential to adsorb on silicate surfaces, and, therefore, suppress  $C_3S$  hydration. By consolidating results pertaining to the three different PCEs, the study advances a simple numerical equation—which unifies the PCE's dosage and architectural parameters into a single numerical value—to assess, in a quantitative manner, a given PCE's potential to suppress  $C_3S$  hydration.

Overall, outcomes of this study provide novel mechanistic insights into the root-cause of decelerating effects of PCE. The discussion provides an improved understanding of how the dosage and architectural parameters of PCE which can readily be characterized using conventional experimental techniques—affect the hydration of C<sub>3</sub>S at early ages. Such knowledge is expected to aid in uncovering the underlying mechanisms that describe the influence of PCE on the hydration of other cementitious phases (e.g., C<sub>3</sub>A), as well as the development of fresh- (e.g., rheology) and hardened-properties (e.g., compressive strength) of cementitious systems.

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**Data availability** The raw/processed data required to reproduce the findings presented in the paper is available upon request.

#### **Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no competing interests.

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