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Exploiting *in-situ* solid-state NMR spectroscopy to probe the early stages of hydration of calcium aluminate cement



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ABSTRACT

We report a high-field *in-situ* solid-state NMR study of the hydration of CaAl₂O₄ (the most important hydraulic phase in calcium aluminate cement), based on time-resolved measurements of solid-state ²⁷Al NMR spectra during the early stages of the reaction. A variant of the CLASSIC NMR methodology, involving alternate recording of direct-excitation and MQMAS ²⁷Al NMR spectra, was used to monitor the ²⁷Al species present in both the solid and liquid phases as a function of time. Our results provide quantitative information on the changes in the relative amounts of ²⁷Al sites with tetrahedral coordination (the anhydrous reactant phase) and octahedral coordination (the hydrated product phases) as a function of time, and reveal significantly different kinetic and mechanistic behaviour of the hydration reaction at the different temperatures (20 °C and 60 °C) studied.

1. Introduction

The application of solid-state NMR spectroscopy to understand fundamental aspects of cement science has a long history [1-3]. Although cements often have complex compositions, solid-state NMR provides key information about the structures of some of the common components, particularly from solid-state ²⁷Al and ²⁹Si NMR measurements. However, while other experimental techniques (including soft X-ray imaging [4] and synchrotron X-ray powder diffraction [5]) have been applied for in-situ studies of hydration processes, previous NMR investigations have been limited to ex-situ studies, involving the examination of samples extracted at different stages during the hydration process and then dehydrated to halt the reaction before measurement of the solid-state NMR data [1,2,6,7]. In part, the focus on ex-situ studies to date has been dictated by technical challenges associated with sealing liquid-containing samples in standard solid-state NMR rotors and then subjecting these samples to magic-angle spinning (MAS) at frequencies of several kHz without leakage of the liquid into the NMR magnet. However, there has been significant progress in recent years in the development of sealing systems for carrying out MAS on liquid-containing samples, which

has facilitated the development of techniques for *in-situ* solid-state NMR studies of crystallization processes from solution [8–15]. In this paper, we report the first application of such *in-situ* NMR techniques to study the hydration of a hydraulic cement, namely calcium aluminate cement, which has a sufficiently high aluminium content to allow ²⁷Al NMR spectra to be recorded with comparatively short data collection times. We note, however, that earlier NMR studies [16–18] examined hydration of silicate materials under similar experimental conditions.

As the ²⁷Al nucleus is quadrupolar (spin I = 5/2), direct-excitation ²⁷Al NMR spectra (recorded using a single ²⁷Al pulse) are typically characterized by broad lines even when the sample is subjected to MAS. While the second-order broadening of the central transition may, in principle, yield information on the electronic configuration around the ²⁷Al nucleus, the line-broadening is often problematic in the study of complex materials or mixtures of solid phases as it can lead to substantial overlap of signals from different ²⁷Al sites. As the line broadening is inversely proportional to the applied magnetic field, it can be advantageous to record ²⁷Al NMR data at high magnetic field. For this reason, the ²⁷Al NMR experiments reported here have been carried out at high magnetic field (20 T; ²⁷Al Larmor frequency, 221.54 MHz) at the U.K.

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National High-Field Solid-State NMR Facility. It is also advantageous to utilize one of the techniques that allow isotropic NMR spectra to be recorded for nuclei of half-integer spin, such as ²⁷Al, and here we use the MQMAS technique [19] in our *in-situ* ²⁷Al NMR studies.

To assess the capability of *in-situ* solid-state ²⁷Al NMR to study cement hydration, we focus on calcium aluminate cement (CAC). Calcium aluminate cements consist primarily of calcium aluminate (CaAl₂O₄) with smaller quantities of mayenite (Ca12Al14O33), calcium dialuminate (CaAl₄O₇) and/or aluminium oxide (Al₂O₃), together with some impurities containing iron and/or silicon. The CAC system was selected for our in-situ solid-state NMR study for three reasons: (i) it is industrially important as a refractory and/or high early strength cement, and as a constituent for specialty mortars (for example, for self-levelling applications and niche applications such as storage of radioactive waste [20-22]), (ii) it has near neutral pH, which minimizes potential hazards for the seals on the NMR rotors used in our in-situ NMR studies, and (iii) it represents a suitable model system due to its simplicity in comparison to other common cements such as Portland cement. Nevertheless, to understand the structural changes that occur as the cement hardens from a freshly mixed slurry to a monolithic solid, time-resolved in-situ analysis is essential.

Different reaction pathways are anticipated in the initial stages of hydration of CAC in different temperature regimes. Below *ca.* 15 °C, reaction of CaAl₂O₄ with water has been observed [23] to produce predominantly the so-called "calcium aluminate decahydrate" (CaAl₂O₄·10H₂O). However, we note that recent work [24] suggests that the stoichiometry of this material actually varies between about 8.3 and 9.3 water molecules per formula unit, as shown in Equation (1):

$$CaAl_2O_4 + y H_2O \rightarrow CaAl_2O_4 \cdot y H_2O (y = 8.3 - 9.3)$$
(1)

Nevertheless, for consistency with previous literature, we refer to this material subsequently as $CaAl_2O_4$ ·10H₂O.

At temperatures in the approximate range [25] $15 \degree C - 30 \degree C$, it has been suggested [22,23] that the main initial product is Ca₂Al₂O₅·8H₂O, possibly with the concomitant formation of CaAl₂O₄·10H₂O (particularly when the temperature is near ambient temperature). Both CaAl₂O₄·10H₂O and Ca₂Al₂O₅·8H₂O are metastable and ultimately undergo dehydration and structural rearrangement to form two stable phases: Ca₃Al₂O₆·6H₂O (tricalcium aluminate hexahydrate, a hydrogarnet type phase) and Al(OH)₃·xH₂O with small values of *x* less than *ca*. 0.2 (a structurally disordered, hydrated form of aluminium hydroxide). Equations (2) and (3) describe this process in a stepwise form:

 $2 \text{ CaAl}_2\text{O}_4 \cdot 10\text{H}_2\text{O} \rightarrow \text{Ca}_2\text{Al}_2\text{O}_5 \cdot 8\text{H}_2\text{O} + 2 \text{ Al}(\text{OH})_3 \cdot x\text{H}_2\text{O} + (9-2x) \text{ H}_2\text{O}$ (2)

$$3 \text{ Ca}_2\text{Al}_2\text{O}_5 \cdot 8\text{H}_2\text{O} \rightarrow 2 \text{ Ca}_3\text{Al}_2\text{O}_6 \cdot 6\text{H}_2\text{O} + 2 \text{ Al}(\text{OH})_3 \cdot x\text{H}_2\text{O} + (9-2x) \text{H}_2\text{O}(3)$$

At sufficiently high temperature [22], hydration of CaAl₂O₄ may lead directly to the formation of Ca₃Al₂O₆· $6H_2O$ and Al(OH)₃· xH_2O , according to Equation (4):

$$3 \operatorname{CaAl_2O_4} + (12+4x) \operatorname{H_2O} \to \operatorname{Ca_3Al_2O_6} + 4 \operatorname{Al(OH)_3} + 4 \operatorname{Al(O$$

In anhydrous CaAl₂O₄, aluminium is present only in sites with approximately tetrahedral coordination, whereas in the hydrate phases, aluminium is present in sites with approximately octahedral coordination (the crystal structures [24,26] of CaAl₂O₄·10H₂O and Ca₃Al₂O₆·6H₂O both have a single octahedrally-coordinated ²⁷Al site). Thus, the isotropic ²⁷Al NMR chemical shifts for the hydrate phases should be clearly distinct from those for anhydrous CaAl₂O₄. However, resolving the ²⁷Al NMR signals for the octahedral sites in the different hydrate phases is more challenging. Indeed, previous *ex-situ* ²⁷Al NMR studies [1,2], in which samples were extracted at different stages during the hydration process and then dehydrated (by treatment with acetone and ether) before recording ²⁷Al NMR spectra, demonstrated the change in coordination of the ²⁷Al sites from tetrahedral to octahedral, but were unable to identify the specific phases produced.

In the recently reported CLASSIC NMR methodology [27], different

types of NMR spectrum are recorded alternately during *in-situ* studies of crystallization from solution, allowing the time-evolution of both the liquid and solid phases to be monitored effectively simultaneously in the same experiment. In the present work, we have implemented a similar strategy involving alternate recording of direct-excitation and MQMAS ²⁷Al NMR spectra during the *in-situ* time-resolved study of hydration of CAC. In the direct-excitation spectra, we expect to observe all ²⁷Al sites in the system, including those in solid and liquid phases. In the MQMAS spectra, on the other hand, we expect to observe signals only from ²⁷Al sites in solid phases for which the multiple-quantum excitation and reconversion pulses are relatively efficient.

2. Experimental Methods

²⁷Al NMR spectra were recorded at 20.0 T on a Bruker AVANCE III spectrometer at the U. K. High-Field (850 MHz) Solid-State NMR Facility (²⁷Al Larmor frequency, 221.5 MHz) with a 4 mm HXY probe (MAS frequency, 10 kHz) and a zirconia rotor with a "high-resolution" insert for liquid-containing samples. The "high-resolution" inserts (also known as "HR-MAS") were purchased from Bruker and used without modification; they are made of Kel-F and consist of a tube into which the sample is placed, followed by sealing with a plug and screw.

Chemical shifts were referenced via a secondary standard of YAG (yttrium aluminium garnet) with the hexa-coordinated site referenced to 0.7 ppm, which corresponds to the standard reference of ²⁷Al in 1 M aqueous Al(NO₃)₃ at 0 ppm. A commercial sample of CAC (Secar 71, Kerneos, France) was used in this work; this material comprises predominantly (ca. 55%) calcium aluminate (CaAl₂O₄) together with calcium dialuminate (CaAl₄O₇) and aluminium oxide (Al₂O₃). For our in-situ ²⁷Al NMR studies, the sample of CAC was mixed with water at a water/ Secar 71 mass ratio of 0.35 (which corresponds to a water/CaAl₂O₄ mass ratio of 0.64). This low ratio was selected as it gives a thick paste, which should minimize the likelihood of phase segregation occurring due to the centrifugation effect of applying MAS to the NMR rotor. The resulting paste was injected into the "high-resolution" insert which was sealed and fitted into the rotor. The rotor was then inserted into the NMR spectrometer, which was already set to the required temperature (20 °C or 60 °C). Sample temperatures were set according to a calibration using lead nitrate [28] and verified in the temperature range of our experiments by measurements on methanol [29-31], taking into consideration the heating effect due to MAS. Data acquisition started within 10 min of the initial mixing of CAC and water.

To record the MQMAS ²⁷Al NMR spectra, a *z*-filtered 3QMAS pulse sequence [32,33] was used with a double-frequency sweep on the 3Q to 1Q conversion pulse [34]. The resulting spectra were sheared using standard Bruker TOPSPIN software. Each direct-excitation spectrum was acquired in 1.1 min (64 scans; recycle delay, 1 s) and each MQMAS spectrum was acquired in 22.2 min (64 t_1 increments with 96 scans and 4 dummy scans per increment; recycle delay, 0.2 s). Thus, the overall time resolution of our *in-situ* NMR study was 23.3 min. Radio-frequency field strengths in the MQMAS experiment were 60 kHz for the excitation pulse, 9 kHz for the selective pulse and 30 kHz for the double-frequency sweep. ¹H decoupling was not applied in any of our experiments.

Isothermal calorimetry experiments were carried out at base temperatures of (20.00 ± 0.02) °C and (60.00 ± 0.02) °C using a TAM Air calorimeter. The fresh cement paste was prepared by external handmixing for 3 min, weighed into an ampoule and immediately placed into the calorimeter to record heat flow for the first 300 h of the hydration reaction. All results are normalized by the total mass of the paste.

3. Results and discussion

3.1. Characterization of the CAC starting material

The direct-excitation 27 Al NMR spectrum of the starting sample of CAC is shown in Fig. 1a. The intense group of peaks between 76 and



Fig. 1. (a) Experimental direct-excitation 27 Al NMR spectrum recorded for the starting sample of CAC [42]. (b) Simulated 27 Al NMR spectrum for a 1.5:1 ratio of CaAl₂O₄ and CaAl₄O₇, calculated using the contour analysis method [43] with line broadening of 0.25 ppm.

84 ppm is due to $CaAl_2O_4$, which has six distinct ²⁷Al environments. The peaks between 50 and 75 ppm are assigned [3] to the two distinct ²⁷Al environments in CaAl₄O₇ and the peaks between 5 and 20 ppm are assigned [35] to Al₂O₃, both of which were also present in the CAC sample (see Experimental Methods). Spinning sidebands due to CaAl₂O₄ are also observed (between 33 and 39 ppm) in Fig. 1a.

Values of isotropic ²⁷Al chemical shifts and ²⁷Al quadrupolar coupling constants for CaAl₂O₄ and CaAl₄O₇ have been reported by Skibsted et al. [3] The simulated ²⁷Al NMR spectrum (Fig. 1b) for a mixture of CaAl₂O₄ and CaAl₄O₇ in 1.5:1 molar ratio, calculated using these isotropic chemical shifts and quadrupolar coupling constants for the ²⁷Al Larmor frequency (221.5 MHz) used in the present work, is in good qualitative agreement with our experimental data (Fig. 1a).

3.2. In-situ 27 Al NMR study of the hydration reaction at 20 °C

Our *in-situ* ²⁷Al NMR experiment on the mixture of the anhydrous CAC sample and water at 20 °C was run for a total of 21 h. The direct-excitation ²⁷Al NMR spectra recorded as a function of time during this experiment are shown in Fig. 2 (only every fifth recorded spectrum is displayed, representing a time interval of 116 min between the spectra shown). First, we note that the contributions due to CaAl₄O₇ and Al₂O₃ remain unchanged as a function of time as these phases do not participate in the hydration reaction. Thus, the changes in the ²⁷Al NMR spectra as a function of time result *only* from the hydration of CaAl₂O₄.

From the start of the hydration process, a new feature appears in the ²⁷Al NMR spectrum between *ca*. 5 ppm and 16 ppm, superimposed on the peak due to Al₂O₃. Clearer insights into the emergence and evolution of the new feature are gained by subtracting the second spectrum recorded [36] (which, in the spectral region from 5 ppm to 16 ppm, contains only the peak due to Al₂O₃) from all subsequent spectra, revealing the growth of the new peak centred at ca. 11 ppm (Fig. 3). Integration of the region between 5 and 16 ppm as a function of time in the data shown in Fig. 3 yields kinetic information on the growth of the new peak and a plot of peak intensity (scaled relative to the peaks for CaAl₂O₄ in the second spectrum) is shown in Fig. 4. It is clear that initial growth of the new peak occurs over the first 5 h of the experiment but the growth is then much slower between 5 h and 16 h. However, after 16 h, the intensity of the new peak increases much more rapidly, signifying a sharp increase in reaction rate, which continues until the end of the experiment. Correspondingly, the intensity of the peaks (between 75 and 86 ppm) due to the CaAl₂O₄ starting material decreases relatively slowly during the first 16 h of the reaction, followed by a much more rapid decrease throughout the remainder of the experiment.



Fig. 2. Direct-excitation 27 Al NMR spectra recorded as a function of time during the reaction of CAC with water at 20 °C: (a) the full spectral region from 0 ppm to 100 ppm and (b) expanded spectra showing the region from 0 ppm to 25 ppm.



Fig. 3. Direct-excitation 27 Al NMR spectra recorded as a function of time during the reaction of CAC with water at 20 °C, following subtraction of the second spectrum recorded. The expanded region from 0 ppm to 25 ppm is shown (as in Fig. 2b).

Our observation that the reaction proceeds in a number of distinct kinetic regimes is consistent with the initial formation of metastable hydrate phases CaAl₂O₄·10H₂O and/or Ca₂Al₂O₅·8H₂O (see Introduction) in the early stages (up to *ca*. 5 hr) of the hydration of CaAl₂O₄, followed by a period of virtual stagnation (between 5 and 16 h) in which very little further formation of hydration products is observed. Significantly, our isothermal calorimetry data for the reaction at 20 °C (Fig. 5) also show evidence for a slow period (representing the first *ca*. 18 hr in this experiment) followed by a period with much more rapid reaction.

Following the slow period of the reaction (up to *ca*. 16 hr in our *in-situ* ²⁷Al NMR study), the metastable hydrate phases CaAl₂O₄·10H₂O and Ca₂Al₂O₅·8H₂O then convert rapidly to the stable products Ca₃Al₂O₆·6H₂O and Al(OH)₃·*x*H₂O [37]. We note that the Al(OH)₃·*x*H₂O produced in this reaction (Equation (3)) is almost always structurally disordered [22] and it is therefore expected to give a significantly broader signal in the ²⁷Al NMR spectrum than the signals for the



Fig. 4. Time-dependence of the intensity of the peaks between 5 and 16 ppm (black; representing the product) and between 75 and 86 ppm (red; representing the reactant) in the direct-excitation 27 Al NMR spectra recorded during reaction of CAC with water at 20 °C. Peak intensities were determined from the spectra after subtracting the second spectrum recorded and are expressed relative to the intensity of the peaks between 75 and 86 ppm for the starting material in the second spectrum.



Fig. 5. Isothermal calorimetry data showing (a) heat flow and (b) cumulative heat flow as a function of time during the reaction of CAC with water at 20 °C (black) and 60 °C (red).

crystalline products of the hydration reaction. For this reason, the new peak that arises at *ca*. 11 ppm in the hydration reaction is not expected to contain any significant contribution from Al(OH)₃·xH₂O.

Conversion of the initially formed metastable hydrates $CaAl_2O_4 \cdot 10H_2O$ and $Ca_2Al_2O_5 \cdot 8H_2O$ to the stable hydrates $Ca_3Al_2O_6 \cdot 6H_2O$ and $Al(OH)_3 \cdot xH_2O$ in the rapid phase of the reaction (after *ca*. 16 hr) is associated with release of water [22,38], as shown in Equations (2) and (3). The water released is available to participate in further hydration of the unreacted $CaAl_2O_4$ particles in parallel with

rapid formation of Ca₃Al₂O₆·6H₂O and Al(OH)₃·*x*H₂O. This mechanism is consistent with the rapid increase in intensity of the peaks for octahedral ²⁷Al sites after *ca*. 16 hr in the direct-excitation ²⁷Al NMR spectra and with an exothermic process (at *ca*. 18 hr) in the isothermal calorimetry data (Fig. 5).

For the reaction at 20 °C, MQMAS ²⁷Al NMR spectra recorded at the beginning of the reaction and after 21 h are shown in Fig. 6. The first spectrum contains two peaks: a broad, high-intensity peak at *ca*. 80 ppm in the direct (1Q) dimension and at *ca*. 83 ppm in the indirect (3Q) dimension, which is attributed to tetrahedral ²⁷Al sites in CaAl₂O₄, and a peak at *ca*. 68 ppm in the direct (1Q) dimension and at *ca*. 78 ppm in the indirect (3Q) dimension, which is attributed to tetrahedral ²⁷Al sites in CaAl₂O₄, and a peak at *ca*. 68 ppm in the direct (1Q) dimension and at *ca*. 78 ppm in the indirect (3Q) dimension, which is attributed to tetrahedral ²⁷Al sites in CaAl₄O₇. After 21 h, a third peak is present, which matches the new peak observed in the direct-excitation ²⁷Al NMR spectra and is attributed to octahedrally coordinated ²⁷Al sites in the hydration products.

3.3. In-situ 27 Al NMR study of the hydration reaction at 60 °C

For the reaction at 60 °C, our in-situ ²⁷Al NMR experiment on a mixture of anhydrous CAC and water was run for a total of 22 h. The direct-excitation ²⁷Al NMR spectra recorded as a function of time after mixing are shown in Fig. 7 (only every fifth recorded spectrum is displayed, with 116 min between the spectra shown). Again, a new peak appears at ca. 11 ppm, superimposed on the peak due to the Al₂O₃ impurity phase, and the intensity of this peak grows rapidly. Integration of the new peak (after subtracting the first spectrum from all subsequent spectra) as a function of time gives insights into the kinetics of the hydration reaction (see Fig. 8, in which peak intensity is scaled relative to the intensity of the peaks for the CaAl₂O₄ starting material in the first spectrum). In Fig. 8, the decrease in the intensity of the peaks due to CaAl₂O₄ (in the region between 75 and 86 ppm) is also shown. The new peak at ca. 11 ppm grows rapidly at the start of the reaction (within the first hour of our experiment) but then slows down to a rate that remains essentially constant for the remainder of the experiment. This behaviour



Fig. 6. *In-situ* MQMAS ²⁷Al NMR spectra (acquisition time, 22 min) recorded during the reaction between CAC and water at 20 °C: (a) the first spectrum recorded and (b) the final spectrum recorded (after 21 h). The indirect dimension was scaled according to the Bruker convention.



Fig. 7. Direct-excitation 27 Al NMR spectra recorded as a function of time during the reaction of CAC with water at 60 °C.



Fig. 8. Time-dependence of the intensity of the peaks between 5 and 16 ppm (black; representing the product) and between 75 and 86 ppm (red; representing the reactant) in the direct-excitation 27 Al NMR spectra recorded during reaction of CAC with water at 60 °C. The peak intensities were determined from the spectra after subtracting the first spectrum recorded and are expressed relative to the intensity of the peaks between 75 and 86 ppm due to the starting material in the first spectrum.

is mirrored by a rapid decrease in the intensity of the peaks due to the $CaAl_2O_4$ starting material within the early period of the reaction, followed by a much slower decrease for the remainder of the experiment.

The first ²⁷Al MQMAS NMR spectrum acquired at 60 °C contains only peaks due to CaAl₂O₄ and CaAl₄O₇. All subsequent MQMAS spectra indicate that these two phases are present, together with a signal corresponding to the new peak observed in the direct-excitation ²⁷Al NMR spectra. Fig. 9 shows the final MQMAS spectrum recorded during the experiment (after 21 h).



Fig. 9. *In-situ* MQMAS ²⁷Al NMR spectrum (acquisition time, 22 min) recorded at the end (21 h) of the reaction between CAC and water at 60 $^{\circ}$ C. The indirect dimension was scaled according to the Bruker convention.

Comparison of Figs. 4 and 8 suggests that there are significant differences in the mechanism of the hydration reaction at 20 °C and 60 °C. At the higher temperature (60 °C), the stable hydrate products $Ca_3Al_2O_6.6H_2O$ and $Al(OH)_3.xH_2O$ are formed rapidly in the early stages of the reaction, in good agreement with isothermal calorimetry data (Fig. 5). It is possible that the metastable hydrate phases $CaAl_2O_4.10H_2O$ and $Ca_2Al_2O_5.8H_2O$ are formed as intermediates at 60 °C, but in this scenario, it is clear that the conversion of these phases to the stable hydrate phases is sufficiently rapid that they are not observed within the time-resolution of our *in-situ* solid-state NMR study (i.e., within 23.3 min).

3.4. Comparison of the hydration reactions at $20 \,^{\circ}C$ and $60 \,^{\circ}C$

At both 20 °C and 60 °C, our direct-excitation ²⁷Al NMR spectra for the hydrated reaction mixture do not contain any measurable contribution from aluminium in the liquid phase at any stage of the reaction, recognizing that solution-state ²⁷Al species would give much narrower peaks than those observed. Fig. 10 shows an overlay of the final ²⁷Al NMR spectra recorded at 20 °C and 60 °C, representing the products present at the end of our experiment at each temperature. Although the peaks for the products are in the same spectral region at each temperature, the peak for the products at 20 °C is somewhat broader and has an observed maximum at slightly higher chemical shift. The ²⁷Al chemical shift observed in each case is fully consistent with ²⁷Al species in sites with approximately octahedral coordination [39]. The fact that the peak due to the products at 20 °C is broader than at 60 °C is consistent with our assertion that the sample at 20 °C contains both metastable hydrate phases CaAl_2O_4 $\cdot 10H_2O$ and Ca_2Al_2O_5 $\cdot 8H_2O$ and the stable hydrate phase $Ca_3Al_2O_6{\cdot}6H_2O$ at the time at which the experiment was stopped, representing a distribution of local structural environments for the ²⁷Al sites. It is clear from Fig. 4 that the reaction to produce the stable hydrate products was still continuing at the end of the experiment at 20 $^\circ\text{C},$ suggesting that some amounts of the metastable hydrate phases were still present.

The significantly different kinetics of the reactions at 20 °C and 60 °C are evident from Figs. 4 and 8. Although temperature was constant throughout the reaction in each case, the rate of reaction varies significantly in different time periods during the experiment. At 20 °C, just over 6% of the CaAl₂O₄ had reacted by the end of the experiment (i.e., 21 h) while, at 60 °C, only about 15% of the CaAl₂O₄ had reacted by the end of the experiment (i.e., 22 h). Assuming in both cases that Ca₃Al₂O₆·6H₂O would be the final product of the reaction, we calculate [40] that the amount of water used in our experiments was more than sufficient to allow the reaction to proceed to completion. Thus, in our experiments at both 20 °C and 60 °C, the reaction did not come close to exhausting the available supply of water. It is likely that the extent of reaction may be limited by the expectation that the hydrate phases are formed as a "coating" on the surfaces of the particles of anhydrous CaAl₂O₄, which hinders the access of water to these particles. This situation does not commonly arise at such low reaction extents during CAC hydration, but it



Fig. 10. Overlay of the peaks in the direct-excitation 27 Al NMR spectra from the new phases produced at 20 °C (black) and at 60 °C (red). In each case, the last spectrum recorded at the end of the *in-situ* 27 Al NMR study is shown (acquisition time, 1.1 min). The spectra are scaled vertically to give similar peak heights.

is possible that the high-frequency (10 kHz) spinning of the NMR rotor in our experiments may influence the accessibility of water to unreacted particles of anhydrous CaAl₂O₄. Unfortunately, very little information is available about the effects of centrifugation on either the process or products of CAC hydration [41].

4. Concluding remarks

We have demonstrated that the hydration reaction of calcium aluminate cement can be monitored successfully by in-situ solid-state ²⁷Al NMR spectroscopy. We emphasize that the CLASSIC NMR methodology used in this study is sensitive to both the liquid and solid phases present during the reaction, and our results demonstrate that no significant amounts of aluminium species are present in the liquid phase at any stage during the reaction. Within the solid phase, our data provide quantitative information on the change in the relative amounts of ²⁷Al sites with tetrahedral coordination (the anhydrous reactant phase) and octahedral coordination (the hydrated product phases) as a function of time. At 20 °C, the reaction rate is initially slow but then accelerates substantially after about 16 h while, at 60 °C, the reaction rate is rapid in the early stages and then slows down considerably. The clear difference in kinetic behaviour at these two temperatures underlines the importance of applying time-resolved in-situ techniques that are sensitive to monitoring the structural changes that occur in the reacting system as a function of time.

The CLASSIC NMR strategy used in the present study involved alternate recording of direct-excitation and MQMAS ²⁷Al NMR spectra during the *in-situ* time-resolved study of hydration of CAC, although (as discussed above) it was found that the MQMAS spectra did not reveal any significant additional information for the reaction system under investigation. Nevertheless, the results from the MQMAS spectra are in full support of our interpretation of the direct-excitation ²⁷Al NMR spectra, and confirm that the observed peaks are definitely from solid phases. Thus, the combined approach of measuring alternating direct-excitation and MQMAS ²⁷Al NMR spectra was still advantageous compared to simply recording only the direct-excitation ²⁷Al NMR spectra.

Finally, although our *in-situ* NMR data were recorded under conditions of rapid sample spinning, it is important to note that the timings of the observed events in the reactions at 20 °C and 60 °C are very similar to those observed in our calorimetry experiments (recorded on a static sample). However, we cannot rule out the possibility that centrifugation caused by sample spinning in the NMR measurements may exert some influence on the reaction system under investigation, for example by altering the distribution of water and CAC within the NMR rotor. Further NMR studies of this reaction system carried out as a function of spinning frequency may provide more detailed insights into this issue.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.

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