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Abstract
A self-degradable alkali-activated cementitious material consisting of a sodium silicate activator, slag, Class C fly ash, and sodium carboxymethyl cellulose (CMC) additive was formulated as one dry mix component, and we evaluated its potential in laboratory for use as a temporary sealing material for Enhanced Geothermal System (EGS) wells. The self-degradation of alkali-activated cementitious material (AACM) occurred, when AACM heated at temperatures of $\geq 200^\circ$C came in contact with water. We interpreted the mechanism of this water-initiated self-degradation as resulting from the in-situ exothermic reactions between the reactants yielded from the dissolution of the non-reacted or partially reacted sodium silicate activator and the thermal degradation of the CMC. The magnitude of self-degradation depended on the CMC content; its effective content in promoting degradation was $\geq 0.7\%$. In contrast, no self-degradation was observed from CMC-modified Class G well cement. For 200°C-autoclaved AACMs without CMC, followed by heating at temperatures up to 300°C, they had a compressive strength ranging from 5982 to 4945 psi, which is $\sim$3.5-fold higher than that of the commercial Class G well cement; the initial- and final- setting times of this AACM slurry at 85°C were $\sim$ 60 and $\sim$ 90 min. Two well-formed crystalline hydration phases, 1.1 nm tobermorite and calcium silicate hydrate (I), were responsible for developing this excellent high compressive strength. Although CMC is an attractive, as a degradation-promoting additive, its addition to both the AACM and the Class G well cement altered some properties of original cementitious materials; among those were an extending their setting times, an increasing their porosity, and lowering their compressive strength. Nevertheless, a 0.7 % CMC-modified AACM as self-degradable cementitious material displayed the following properties before its breakdown by water; $\sim$ 120 min initial- and $\sim$180 min final-setting times at 85°C, and 1825 to 1375 psi compressive strength with 51.2 to 55.0 % porosity up to 300°C.
Introduction

A critical operation in assembling and constructing the enhanced geothermal system (EGS) is to create the hydrothermal reservoir in an impermeable rock stratum at temperatures $\geq 200^\circ C$, located at $\sim 3$-$10$ km below the ground surface. In this operation, water is pumped down from injection well to stimulate the hot rock stratum. This hydraulic stimulation initiates the opening of existing fractures [1,2]. The continued pumping of water acting as stimulation fluid causes the fractures to open after which the rock faces then shift due to existing shear stress. When the fracture close back up after stimulation, the irregularities in the rock prevent them from closing completely which results in the formation of a permeable fracture flow network. After forming the reservoir, a production well is installed within the fracture’s network. Then, the heat transmission fluid as the working fluid from the injection well is transported to the production well through the hot fracture zones; thereafter, it circulates between the injection- and production-wells.

During such wellbore construction, operators pay considerable attention to the pre-existing fractures, and pressure-generated ones made in the underground foundation during drilling and logging. These fractures in terms of lost circulation zones often cause the wastage of a substantial amount of the circulated water-based drilling fluid or mud. To deal with this problem, operators must seal or plug the lost circulation zones with appropriate materials to avoid depleting the drilling fluid and loosing mud. Once this problem is resolved, the drilling operation can resume and continue until the wellbore structure is completed. Thereafter, the hydraulic –stimulation process begins. Next the inevitable concern is the fact that all sealing materials that were used to plug the fractures present in hydrothermal reservoir’s rock stratum must be disintegrated by highly pressured water to reopen the sealed fractures and to promote the propagation of reopened fractures.

Presently, Ordinary Portland Cement (OPC) not only is used as a well-casing cementing material, but also often is adapted as the binder in the sealing systems including bridging additives like fibrous materials [3,4]. This fact suggested that in developing the ideal
sealing binders, their potential value for use as well-casing cements also should be considered. Earlier, when the OPC-based cementitious materials were employed widely as geothermal well-casing cement, which serves in supporting mechanically the metal casing-pipe and in protecting it against corrosion, one serious drawback was their limited resistance to the hot acidic environment created by the combination of concentrated dihydrogen sulfide (H₂S) and carbon dioxide (CO₂) in geothermal wells. As well documented [5,6], the capacity of OPCs to withstand acid was very poor, and they suffered from severe acid erosion. To deal with this problem, acid-resistant cements had to be developed.

In response to this need, researchers at Brookhaven National Laboratory (BNL) formulated alkali-activated cementitious materials (AACMs) that resisted acid much better than did the OPCs; this work was part of DOE’s earlier geothermal program aimed at developing alternative cost-effective acid-resistant geothermal well cements [7,8]. The AACMs were prepared hydrothermally using two major components: One was inexpensive industrial by-products with pozzolanic properties, such as the slag from steel industries and the fly ashes from coal-combustion power plants; the other was the sodium silicate along with various mol. ratios of Na₂O/SiO₂ as the alkali activators that initiate the pozzolanic reactions. Two factors contributed to minimizing acid erosion: One was the self-repairing property of the cement itself; the other was the anti-acid zeolite phase formed by the interactions between the mullite in fly ash and the Na ions liberated from the sodium silicate activator. From this work, we concluded that these economical AACMs have a high potential as acid-resistant geothermal well cements at temperatures up to 200°C.

In a standpoint of the self-degradation of cementitious materials, the biodegradable biopolymers including the starch, cellulose acetate, gelatin, and poly(L-lactic acid) in the form of powder, microsphere, and fiber, are currently used as additives, which promote the partial biological degradation of biocompatible bone cements [9-12]. When these additives in the bone cements came in contact with body fluids, they are degraded to create high porosity of cements, thereby providing inter-connective channels for the bone
tissue’s growth and facilitating the resorption of cement. In addition, our interest in biopolymers was their intriguing mechanism of thermal degradation. The cellulose and cellulose-related compounds are degraded thermally in air at temperatures around 200°C; yielding acetic acid as by-products [13-15]. This information inspired us to investigate the potential of cellulose compounds as thermal degradable additives to promote the self-degradation of temporary cementitious sealing materials at temperatures $\geq 200^\circ$C after water has penetrated through the sealer.

Based upon this concept, our focus in this study centered on formulating AACMs, in particular, slag/Class C fly ash combined system, as the binder in temporary sealing material, and investigating the ability of sodium carboxymethyl cellulose (CMC) as thermal degradable additive to promote the disintegration of AACM-based sealer. CMC and cellulose-related compounds were not new materials in drilling industries. These compounds frequently are used as additives in water-based drilling mud to reduce fluid loss in mud, to confer water-holding properties on mud, and to assure its appropriate rheology at elevated temperature [16-20].

In obtaining this information, this study had the following four major objectives: First objective was to optimize the formula for blending the alkali-activated slag/Class C fly ash dry components, using data on its setting time, compressive strength, and identification of crystalline phases responsible for strengthening the sealant. The second objective focused on the thermal decomposition of CMC additive, and the chemistry of decomposed CMC. The third one was measuring the changes in setting time, porosity, and compressive strength of optimized AACM as a function of CMC content, and also analyzing the chemical behavior of CMC in AACM in different thermal environments. The final one was to visualize the self-degradation of the CMC-modified AACM sealer and to determine in-situ exothermic energy evolved in the sealer while water penetrated through them. The integration of all information described above was directed toward understanding the self-degradation mechanism.
**Experimental Procedure**

Two industrial by-products with pozzolanic properties, granulated blast-furnace slag under trade name “New Cem,” and Class C fly ash, were used as the hydraulic pozzolana cement. The slag was supplied from Lafarge North America, and fly ash was obtained from Boral Material Technologies, Inc. Table 1 lists their chemical constitutions. A sodium silicate granular powder under trade name “Metos Bead 2048,” supplied by The PQ Corporation, was used as the alkali activator of these pozzolana cements, and its chemical composition was 50.5 mol. wt % Na₂O and 46.6 mol. wt % SiO₂. The formula of the dry pozzolana cements employed in this test had slag/Class C fly ash ratios of 40/60, 20/80, and 0/100 by weight. Sodium silicate powder activator of 4 % by the total weight of pozzolana cement was added to prepare the dry mix cementitious reactant. For comparison purpose, the commercial Class G well cement was used as control.

The sodium carboxymethyl cellulose (CMC) was produced by etherification of cellulose from renewable resources, like wood, and had chemical structure shown in Figure 1. CMC under the production name “Walocel CRT 30 PA,” supplied by Dow Chemical Corp., was used as the thermal degradable additive to promote the disintegration of the AACM-based sealers; it dissolves in water, forming an anionic polymer. A 0.4, 0.7, and 1.0 % CMC by total weight of dry slag/Class C fly ash/activator mixtures was incorporated into AACMs.

AACM slurries with and without CMC were prepared by adding an appropriate amount of water to the dry mix cementitious component, and then left them in an atmospheric environment until the cement set. Adding the CMC led to the entrainment of air into slurries, reflecting the formation of lightweight slurries. Afterward, the room temperature-set AACM samples were exposed in an autoclave at 200°C for 5 hours; some autoclaved samples then were heated for 24 hours in over at 200°, 250°, and 300°C.

**Measurements**

The initial- and final-setting times of CMC-incorporated and unincorporated AACM sealing slurries at 85°C were determined in accordance with modified ASTM C 191-92.
In it, a slurry-filled conical mold was placed in screw-topped round glass jar to avoid any evaporation of water from slurry during heating. The slurry was examined every 30 minutes to determine its setting time. The porosity of autoclaved and heated AACMs was measured by helium pycnometry. The specimens with size, 35 mm diameter x 70 mm length were used for porosity and compressive strength testing. X-ray powder diffraction (XRD) was used to identify the phase composition of the autoclaved- and heated-samples.

With a Type K thermometer, in conjunction with temperature data logger, we monitored the heat energy generated in the AACM slurries during the induction period of its hydration reactions, and also to determine the exothermic energy evolved in a self-degrading process of the cured AACM specimens after their contact with water.

Differential scanning calorimetry (DSC) and Fourier transform infrared spectroscopy (FT-IR) give us information on the thermal decomposition and chemistry of CMC; further, we adapted FT-IR to reveal the chemical behavior of the decomposed CMC in AACM. In preparing the film sample of CMC, 1.0 g “as-received” CMC powder was dissolved in 200 g deionized water (D.I.). A certain amount of CMC solution was poured into a cell culture dish, and heated at 85°C for 24 hours. The resulting CMC film then was exposed for 24 hours in an oven at temperatures of 200°, 250°, and 300°C.

**Results and Discussion**

**Properties of Slag/Class C fly ash Blending System**

Our first experiment was directed towards gaining data on the setting time and compressive strength of AACM samples made with slag (S)/Class C fly ash (C) ratios of 40/60, 20/80, and 0/100. Table 2 shows the changes in water content, expressed as the ratio of water (w) to dry cement (c), as a function of the S/C ratio used in preparing the AACM slurries; it includes their initial- and final-setting times at 85°C. The water content in these AACM systems was adjusted to attain a consistency similar to that of Class G well cement slurries made with the w/c ratio of 0.39. As evident, the w/c ratio decreased with the replacement of some portion of slag by Class C fly ash. The w/c ratio
of slurry made with 0/100 S/C ratio without slag was ~ 20% and ~ 30% lower than that of
the 40/60 S/C ratio and the Class G well cement slurries, respectively. For the latter, no
initial set was apparent after exposure for 60 min in an oven at 85°C, but extending the
exposure time to 90 min resulted in its final set, demonstrating that both its initial- and
final-setting occurred within 30 min. In contrast, all AACMs had at least 30-min interval
between the initial and final settings. The elapsed time for AACM’s initial setting
depended on the S/C ratios; with 40/60 S/C ratio, it was ~ 30 min, corresponding to a 60
min shorter than that of the Class G slurries. This initial setting time extended to ~ 60 min
when 20% of slag was replaced by Class C fly ash, beyond this, as in the 0/100 S/C ratio,
the fly ash was ineffective in further extending setting time. As a result, it appeared that
the slag acted to accelerate the setting of slurries.

Figure 2 compares the compressive strength of the specimens made with 40/60, 20/80,
and 0/100 S/C ratios after exposure in four different thermal conditions, as described
under Experimental Procedure. For comparison purpose, the Class G well cement also
was tested under the same conditions. After autoclaving at 200°C, the samples with a
40/60 S/C ratio had a compressive strength of 5144 psi; those made with the 20/80 ratio
displayed a further ~ 16 % improvement to 5981 psi. The 0/100 ratio without slag was
not as effective in improving strength as that of 40/60 and 20/80 ratios, strongly
demonstrating that incorporating an appropriate amount of slag into Class C fly ash was
essential in attaining the best strength in autoclaved AACMs. In contrast, the effect of
autoclaving on the development of strength in commercial Class G well cement was poor,
reflecting ~ 3.8- and ~ 2.4-fold lower than that of 20/80 and 0/100 ratios, respectively.
Heating the autoclaved AACMs at 200°, 250°, and 300°C engendered a decline in
compressive strength of all them that was directly related to increasing temperature. For
instance, the strength of the autoclaved 20/80 ratio fell ~17% to 4945 psi after heating at
300°C. There was no significant change in strength for the Class G cement in this
temperature range.

From this result, the 20/80 S/C ratio that possessed the highest compressive strength and
initial setting of ~ 60 min was selected as matrix of sealing materials. Figure 3 depicts the
XRD tracings for autoclaved 20/80 ratio specimens before and after heating at 300°C, and also it includes “as-received” slag and Class C fly ash as the starting materials. For slag, XRD pattern (a) revealed the typical amorphous feature of CaO-Al₂O₃-MgO-SiO₂ glass structure. The pattern (b) for Class C fly ash consisted of four crystalline phases, quartz (SiO₂), calcium sulfate anhydrite (CaSO₄), tricalcium aluminate (3CaO·Al₂O₃, C₃A), and lime (CaO). After autoclaving the sodium silicate-activated 20/80 S/C ratio specimen at 200°C, two hydrated reaction products were formed in the cementitious body through the hydrothermal-catalyzed pozzolanic reactions (see pattern c): One was 1.1 nm tobermorite [Ca₃(OH)₂Si₆O₁₆·4H₂O] ; the other was related to calcium silicate hydrate (I) [CaO·SiO₂·H₂O, C-S-H (I)], while some non-reacted phases, such as C₃A and quartz attributed to the Class C fly ash remained. Neither calcium sulfate nor lime was detected in this pattern, seemingly suggesting that these calcium compounds were transferred to the reaction products during autoclaving. To obtain more information on the crystalline reaction products, we surveyed the phase composition of autoclaved 0/100 S/C ratio without slag by XRD (Figure 4). Compared with the composition of the autoclaved 20/80 ratio, the slag-free AACM comprised poorly developed 1.1 nm tobermorite and C-S-H (I) phases, reflecting the weak intensity of d-spacing lines related to these phases. Thus, the addition of some slag to Class C fly ash in AACM system was responsible for developing well-formed tobermorite and C-S-H (I) phases, both of which improve the strength of AACM. Upon heating at 300°C, as shown in Fig. 3, its pattern (d) revealed no changes in the composition of well-crystallized phases, compared with that of autoclaved one, suggesting that these phases formed by autoclaving withstand 300°C heating. Nevertheless, both well-formed tobermorite and C-S-H (I) phases were responsible for strengthening the 20/80 ratio AACM.

Next, our focus shifted to investigating the properties of CMC additive and assessing the potential of incorporating it into the 20/80 ratio AACM as a self-degradable sealer.

**CMC Additive**

Figure 5 gives the DSC curve of the CMC at temperatures from 100°C to 450°C in nitrogen; it encompasses two endothermic-transition peaks, \( T_{p1} \) and \( T_{p2} \), at 270°C and
304°C. Since the endothermic peaks reflect the thermal decomposition of the samples, CMC seemingly has a two-stage decomposition process. The onset of the first stage began at 165°C, denoted as $T_{o1}$ and ended at 295°C, marked as $T_{o2}$. The second stage occurred between 295°C and ended at 322°C as $T_e$. To support this information, we determined the exothermic decomposition energy evolved on each stage of CMC’s thermal decomposition from the closed areas of curves with the baseline. The values for the first- and second-stages were 510.8 and 53.8 J/g, respectively, meaning that the most of CMC’s decomposition took place at temperatures between 165°C and 295°C.

Figure 6 illustrates FT-IR spectra in the wavenumber region, 3500-700 cm$^{-1}$, of CMC non-heated and heated at 200°C, 250°C, and 300°C for 24 hours. Based upon the chemical structure of CMC in Figure 1 in conjunction with literature survey [21-23], the spectrum (a) of the 80°C-made CMC film included several prominent absorption bands at 2925 and 2864 cm$^{-1}$ associated with C-H anti-symmetric and symmetric stretching vibrations, at 1603 cm$^{-1}$ belonged to aromatic C=C stretching mode, at 1576 cm$^{-1}$ shoulder attributed to C-O anti-symmetric stretching in the carboxylate ion group, -COO$^-$, at 1400 cm$^{-1}$ due to -CH$_2$ scissoring overlapped with C-O symmetric stretching in -COO$^-$, at 1329 cm$^{-1}$ related to both -CH$_2$ scissoring and C-OH bending, at 1157 cm$^{-1}$ referred to C-O-C stretching in the ether group, and both at 1057 and 1020 cm$^{-1}$ assignable to C-O stretching in >CH-O-CH$_2$- linkage. Compared with this, FT-IR spectrum (b) of the 200°C-heated CMC film was characterized by declining the band intensity at 1329, 1057, and 1020 cm$^{-1}$, and disappearing the band at 1157 cm$^{-1}$, demonstrating that heating the CMC at 200°C entailed the breakage and rupture of several linkages, such as C-OH, C-O-C, and >CH-O-CH$_2$- within CMC’s molecular structure. Further increasing temperature to 250°C led to the incorporation of five new bands at 1685, 1457, 879, 835, and 780 cm$^{-1}$ into the spectrum (c). The possible contributors to these new bands at 1685 and 1457 cm$^{-1}$ were the carbonyl, C=O, in ketone species formed by intra-ring dehydration [24] and a carbonated compound like sodium carbonate, Na$_2$CO$_3$ [25], respectively. The other new bands emerging in low wavenumber region < 900 cm$^{-1}$ might be associated with the organic fragments yielded by thermal degradation of CMC. This spectrum also strongly represented that the band at
1576 cm⁻¹ attributed to –COO⁻ ionic group had become one of the major peaks, while the aromatic- and >CH-O-CH₂- linkage-related bands had vanished. Thus, the thermal decomposition of CMC at 250°C seemingly generates many –COO⁻ groups that are converted into organic acid in an aqueous media. Several investigators [13,15] reported that this thermal decomposition generated volatile decomposition products, such as CO, CO₂, CH₄, water, and acetic acid. Among those, the major decomposition product at 250°C was acetic acid, which occupies more than 40 % of its total amount. This information strongly supported the likelihood of –COO⁻ → organic acid transformation as well as the formation of sodium carbonate from the interactions between two decomposition by-products of CMC, sodium oxide and CO₂. At 300°C, the spectrum (d) revealed the incorporation of more C=O group and Na carbonate into the thermally degraded CMC, as reflected in the pronounced growth of bands at 1685 and 1457 cm⁻¹ coexisting with the new band at 849 cm⁻¹ attributed to carbonate.

**CMC-incorporated 20/80 S/C Ratio AACM**

Figure 7 shows the changes in temperature of the 20/80 S/C ratio AACM and Class G cement slurries within 530 sec after mixing with water. Shortly after mixing dry AACM with water, its temperature rose from 21°C to 24°C, and continuously increased until it leveled at 27.5°C after 290 sec. This rapid rise was due to the dissolution of the sodium silicate activator in an aqueous media. As expected, Class G cement slurry doesn’t generate any heat in such a short hydration period.

Table 3 lists the water-to-cement (w/c) ratios and the initial- and final-setting times at 85°C for Class G well cement and 20/80 S/C ratio AACM modified with 0.4, 0.7, and 1.0 wt% CMC. In both the cement systems, the w/c ratio increased with an increasing CMC content, from 0.39 at 0 % CMC to 0.62 at 1.0 % CMC for Class G cement, and from 0.31 at 0 % CMC to 0.52 at 1.0 % CMC for AACM. Adding the CMC to the Class G cement remarkably prolonged its setting time; in fact, the value of initial setting time without CMC was extended by nearly 210 min to ~300 min after incorporating 0.4 % CMC. Correspondingly, adding more than 0.4 % CMC considerably delayed the hydration of cement so there was no evidence of initial setting time within 360 min. Undoubtedly,
CMC acted as set retarder of Class G cement. In contrast, CMC was not as effective in retarding the setting of AACM. All 20/80 S/C ratio AACM modified with CMC were set for curing period up to ~ 210 min. With 0.4 % CMC, both the initial- and final-setting times of CMC-free AACM were prolonged by ~ 30 min to ~ 90 and ~ 120 min. Further increasing amount to 0.7 % led to the extension of another 30 and 60 min of initial- and final-setting times over that of 0.4 %. Nevertheless, at the room temperature of ~ 25ºC, the final setting of both the Class G and AACM specimens was observed within 30 hours after mixing with water.

Figures 8 and 9 give the changes in compressive strength of AACM and Class G cement exposed in the four different thermal environments described earlier as a function of CMC content. For AACM, the strength for all specimens exposed to all different environments fell with increasing CMC content; in particular, the addition of the first 0.4 % CMC resulted in a remarkable decline in strength, ranging from 42-60%, compared with that of CMC-free specimens; beyond this content, strength fell monotonously. However, despite adding a high CMC content of 1%, all specimens, except for those heated at 300ºC, retained a compressive strength of more than 1000 psi. A similar trend was observed from Class G cement, namely, the strength decreased with an increasing CMC content. In fact, the compressive strength values, ranging from 1594 to 1475 psi, for all CMC-free specimens fell to less than 1000 psi when ≥0.7% CMC was added.

Figures 10 and 11 plot the porosity vs. CMC content for the AACM and Class G cement after expose to different heat treatment conditions. For AACM, after drying autoclaved specimens without CMC for 24 hours at 85ºC, their porosity was 12.5 %. This value increased by ~2.3 times to ~ 29.0 % when the specimens were heated in the temperature range from 200º to 300ºC. As described earlier, XRD study revealed that the crystalline phase composition of autoclaved AACM after heating at 300ºC closely resembled that of non-heated one. Thus, the increased porosity after heat treatment is more likely to be associated with the elimination of a certain amount of free water linked to the cementitious matrix, rather than the dehydration of crystalline hydration products as the hydrothermal reaction products. In heated CMC-containing AACM systems, CMC
dramatically impacted in creating the porous cementitious structure. Adding 0.4% CMC resulted in the formation of AACM having the porosity > 40%. Further, raising the temperature promoted the incorporation of more voids into the AACM. Both autoclaved and heated AACMs amended with 1% CMC displayed a porosity of more than 50%. For Class G cement, CMC-free autoclaved cement had a porosity of 35.5%, and this value rose to ~40% by heating at 200°-300°C. Like CMC-modified AACM, adding CMC to Class G caused the incorporation of more voids into cement structure, while the increase in heating temperature engendered the incorporation of additional voids.

As mentioned in the Experiment Procedure and discussed in Table 3, CMC was responsible for an air entrainment and an increase in water/cement ratio of AACM that assured the constant consistency of cement slurries. Thus, there were two possible reasons for the creation of porous structure: One was the incorporation of air into the AACM’s slurries; the other was the increased water content of the slurries. To elucidate fully the role of CMC in increasing the porosity, we analyzed 1.0% CMC-modified 20/80 S/C ratio specimens by FT-IR. All specimens were dried in an oven at 85°C for 24 hours before analyzing them. The “as-received” slag and Class C fly ash were also used as references. Figure 12 illustrates the FT-IR spectra of these references; for slag, the spectrum had three prominent bands at ~2923 cm⁻¹, ~1620 cm⁻¹, and ~1430 cm⁻¹, and one wide band in the range of 1050 to 820 cm⁻¹. The first two bands are associated with water adsorbed into cementitious body, while the third band is due to the carbonated compounds. In contrast, no carbonation product was found in Class C fly ash. Figure 13 compares the spectral features of specimens made under the room temperature for 24 hours and under four different conditions. The spectrum (a) of the room temperature-cured specimen, followed by heating at 80°C had CMC-related bands at 1597, 1405, and 1354 cm⁻¹ in conjunction with that from carbonates at 1450 cm⁻¹, and from water at 2923, 2853, and 1627 cm⁻¹. Similar spectral feature, except for the appearance of carbonate-related band at 1420 cm⁻¹, was obtained from specimen autoclaved at 200°C for 5 hours (b), strongly suggesting that CMC didn’t decompose in such a lengthy hydrothermal environment. In comparison, the features of 200°C-heated specimen (c) was different; in particular, the elimination of all CMC-related bands and the marked growth of band at
1420 cm$^{-1}$. As discussed in the DSC study of CMC, the thermal degradation of CMC begun at 165°C, subsequently, causing the emission of CO$_2$. Hence, it is reasonable to consider that the in-situ dry carbonation reactions between the CO$_2$ and alkaline earth metals, like Ca from slag and Class C fly ash or alkaline metals like Na from CMC occurred in cementitious bodies during heating, and lead to the formation of carbonation products, such as CaCO$_2$ and Na$_2$CO$_3$. At 250° (d) and 300°C (e), their spectral features closely resembled to that of 200°C. Accordingly, we conclude that the generation of gasses phase emitted by thermal degradation of CMC was the other factor governing the increase in porosity of AACMs.

**Self-degradation and Its Mechanism**

We questioned whether CMC-modified AACM really degrades after contacting with water. In respond to this question, two different tests were conducted to visualize its self-degradation: One test was directly to immerse hot 20/80 S/C ratio AACM and Class G cement modified and unmodified with CMC into water at 25°C; the other was to impregnate water into the air-free CMC-modified and unmodified 20/80 ratio AACM and Class G cement specimens. The detailed test procedures were as follows; for the first test, 1) the specimens was autoclaved at 200°C for 5 hours, and then heating at 200°C for 24 hours, 2) shortly after removing hot specimens form an oven, they were immersed in the water at 25°C, and 3) visual observation allowed us to evaluate the magnitude of its self-degradation. For the second test, 1) the specimens were autoclaved at 200°C for 5 hours, and then heated at 200°C for 24 hours, 2) the specimens were cooled for 24 hours at room temperature, 3) thermocouple-embedded specimens were placed in a vacuum chamber at -1000 kPa for 10 min to eliminate any air present in the specimens, 4) air-free voids in the vacuumed specimens were filled with water (water/specimens volume ratio= 17/1), while determining the generation of in-situ exothermic heat by temperature data logger, and 5) visual observation was made to evaluate the magnitude of self-degradation.

For the first test, figure 14 shows the appearance of 200°C-heated 20/80 ratio AACMs containing 0, 0.4, 0.7, and 1.0% CMC after the immersing them in water at 25°C. Shortly after immersion, we observed the development of numerous cracks in all CMC-modified
AACMs. Furthermore, the propagation of cracks with elapsed immersion time led to their disintegration. We visually observed that AACMs modified with CMC of ≥ 0.7 % seem to generate a large number of wider crack, compared to specimens with 0.4 % CMC, emphasizing that the higher concentration of CMC promoted more self-degradation in AACM. In contrast, CMC-free AACM showed no signs of degradation. We noted a similar magnitude of degradation in CMC-modified AACMs heated at 250°C and 300°C (data not shown).

In contrast, incorporating CMC into the Class G cement didn’t affect degradation; there was no development of any cracks. This information strongly demonstrated that CMC conferred potential self-degrading properties on AACM, but not on commercial Class G well cement. A possible interpretation for this fact was that when the water was permeated into AACM containing thermally decomposed CMC, it served in disintegrating the hard cementitious structure possessing a compressive strength of more than 2000 psi. Conceivably, such disintegration by water may be due to the generation of water-catalyzed energy in AACM. If this assumption is valid, two resources were involved, one from AACM, and the other from the decomposed CMC.

To verify this assumption, next, our attention turned to the finding from the second test, i.e., determining the exothermic heat energy generated in AACM by water penetration. Plotting the relation curve between the elapse time and exothermic temperature for 0.7 % CMC-modified AACM in Figure 15 shows that ~ 130 sec after impregnating the vacuumed specimen with water, its internal temperature rapidly rose from 22.2°C to a peak of 41.5°C. The total elapsed time from the onset to the maximum temperature was only 110 sec. We observed the development of numerous cracks in the water-impregnated specimens along with the increase in their exothermic temperature. Hence, the penetration of water through the specimen led to the evolution of in-situ exothermic heat promoting the self-degradation of the sealer. Figure 16 plots the peak temperature of heat energy generated in the AACM and Class G cement specimens containing 0, 0.4, 0.7, and 1.0 % CMC. For AACM, although the specimen does not have any CMC, the uptake of water generated some degree of in-situ heat; in fact, specimen’s original
temperature of 22.2°C rose to 32.6°C after impregnation. As we noted earlier, the dissolution of the sodium silicate activator in an aqueous media evolves heat. Correspondingly, this rise in temperature primarily was caused by the re-dissolution or dissolution of partially dissolved or non-dissolved sodium silicate activator remained in the 200°C-heated AACM after water uptake. Of particular interest was the further increase in internally generated heat when CMC was added to it. With 0.4 % CMC, the exothermic temperature rose by 8°C to 40°C. Incorporating more CMC further enhanced its internal temperature. As is seen in Figure 16, the magnitude of self-degradation depended on the exothermic heat generated, which corresponded to CMC content; a high exothermic temperature severely decomposed the sealing material.

In contrast, no internal heat energy was produced in Class G cements modified with CMC; accordingly, the specimens remained intact in terms of non-self degradation.

Integrating all these data, we propose the self-degradation mechanism of CMC-modified AACM depicted in Figure 17, as follows. Although our study provide limited information on the hydrothermal degradation of CMC, no degradation of CMC additive was found in AACM body after autoclaving at 200°C for 5 hours. In the first step of self-degradation routs, when the autoclaved sealing material was heated at temperatures of ≥ 200°C, the CMC in the AACM thermally decomposed, yielding five reactants, CO₂, CO, CH₄, CH₃COOH, and Na₂CO₃. In the second step, the penetration of water through AACM triggered the dissolution of non-reacted or partially reacted sodium silicate activator to release sodium hydroxide, NaOH, as hydrolysate. The sodium silicate next reacted with ionic carbonic acid, CO₃²⁻ and 2H⁺, known as wet carbonation to form sodium bicarbonate, NaHCO₃. In a final step, sodium bicarbonate favorably reacted with acetic acid, CH₃COOH, from CMC to yield three reaction products, sodium acetate, carbon dioxide, and water. This exothermic reaction is very similar to that between baking soda and vinegar well known as the “volcanic reaction.” Thus, the emission of carbon dioxide evolved from this reaction might expand the volume of cementitious structure, promoting its in-situ degradation. On the other hand, the sodium carbonate derived from thermal decomposition of CMC might undergo wet carbonation, Na₂CO₃ +
CO$_3^{2-}$ + 2H$^+$ → 2NaHCO$_3$. However, the total amount of NaHCO$_3$ yielded from this wet carbonation, if any, was very small, and insufficient to degrade the cementitious structure by its following reaction with CH$_3$COOH. The other possibility for promoting the self-degradation was the acid-base exothermic reactions between sodium hydroxide and acetic acid.

**Conclusion**

An alkali-activated cementitious material (AACM) consisting of 3 wt% sodium silicate (Na$_2$O/SiO$_2$ = 1.08 mol. ratio), 14.7 wt% slag, 59.3 wt% Class C fly ash, and 23 wt% water was selected from among several candidate formulas, and its properties were compared with commercial Class G well cement to evaluate its a potential as matrix of temporary, self-degradable cementitious sealers. At 85°C, AACM slurry had initial- and final-setting times of ~ 60 min and ~ 90 min, respectively, and developed a compressive strength of 5981 psi after autoclaving for 5 hours at 200°C. This value was 3.8-fold higher than that of Class G well cement autoclaved at the same temperature. Two well-formed crystalline phases, 1.1 nm tobermorite and calcium silicate hydrate (I), were responsible for developing this excellent compressive strength. Exposing autoclaved AACM in an air-oven at temperatures from 200° to 300°C caused a slight decline in its compressive strength with an increasing temperature. However, at 300°C, its compressive strength still remained at nearly 5000 psi, and these crystalline phases formed in autoclaved AACM were intact.

Sodium carboxymethyl cellulose (CMC) powder was employed as a self-degradation promoting additive of AACM. A 0.4, 0.7, and 1.0 % CMC was incorporated into dry mix AACM. The thermal decomposition of CMC began at 165°C, and was completed at 322°C. At ≥ 200°C, its decomposition generated five by-products, CO$_2$, CO, CH$_4$, CH$_3$COOH, and Na$_2$CO$_3$. However, the CMC in AACM didn’t decompose during autoclaving for 5 hours at 200°, suggesting that the thermal behavior of CMC in a hydrothermal environment differed from that in a dry-heat environment. Adding CMC changed the properties of AACM and Class G well cement; it prolonged their initial- and final-setting times, reduced compressive strength, and increased their porosity.
case, CMC acted as a setting retarder; for AACM, 0.4 % CMC extended its initial- and final-setting time to ~ 90 min from ~ 60 min and to ~120 min from ~ 90 min, respectively. For Class G well cement, CMC was unsuitable as an additive; with 0.4 % CMC, the cement slurry didn’t set within 360 min at 85°C. Regarding the increase in porosity, three factors operated in creating porous structure in AACM and led to a decline in compressive strength: First was the entrainment of air into slurries by CMC, second was an increase in water/cement ratio as a function of the CMC content; and third was the greater emission of gas by-products, such as CO₂, CO, CH₄ with an increasing CMC content.

When CMC-incorporated AACM heated at temperatures ≥ 200°C came in contact with water, the water penetrating through it played pivotal role in promoting the self-degradation of AACM. There were two important roles: One was the dissolution of non-reacted and partially reacted sodium silicate activator remaining in the AACM, reflecting the release of sodium hydroxide as reactant; the other was the formation of ionic carbonic acid from wet carbonation of CO₂ deduced from thermal degradation of CMC. The sodium hydroxide released reacted directly with ionic carbonic acid to form sodium bicarbonate. Then, the chemical affinity of sodium bicarbonate with CH₃COOH from the decomposed CMC led to the evolution of CO₂ as reaction product. In-situ exothermic energy was generated throughout this process, including dissolution, interactions, and evolution. Nevertheless, we consider that this in-situ CO₂ gas evolution led to volume expansion of AACM, thereby resulting in the breakage and disintegration of AACM possessing a compressive strength of more than 2000 psi. In contrast, no self-degradation was noted from CMC-containing Class G well cement.
References


Table 1. Chemical composition of Ground Granulated Blastfurnace Slag and Class C fly ash.

<table>
<thead>
<tr>
<th></th>
<th>CaO, wt%</th>
<th>SiO₂, wt%</th>
<th>Al₂O₃, wt%</th>
<th>MgO, wt%</th>
<th>Fe₂O₃, wt%</th>
<th>TiO₂, wt%</th>
<th>Na₂O, wt%</th>
<th>K₂O, wt%</th>
<th>SO₃, wt%</th>
<th>Loss in ignition for C, wt%</th>
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<tbody>
<tr>
<td>Slag</td>
<td>38.5</td>
<td>35.2</td>
<td>12.6</td>
<td>10.6</td>
<td>1.1</td>
<td>0.4</td>
<td>-</td>
<td>-</td>
<td>0.1</td>
<td>1.5</td>
</tr>
<tr>
<td>Class C fly ash</td>
<td>25.6</td>
<td>36.2</td>
<td>19.1</td>
<td>5.4</td>
<td>6.7</td>
<td>-</td>
<td>1.7</td>
<td>0.5</td>
<td>1.7</td>
<td>3.1</td>
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</table>

Table 2. Water/cement ratio, and initial and final setting times at 85°C for slurries made with various S/C ratios and Class G well cement.

<table>
<thead>
<tr>
<th>Water/cement, w/c, ratio</th>
<th>Initial setting time at 85°C, min</th>
<th>Final setting time at 85°C, min</th>
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<tbody>
<tr>
<td>Class G well cement</td>
<td>0.39</td>
<td>~90</td>
</tr>
<tr>
<td>40/60 S/C</td>
<td>0.34</td>
<td>~30</td>
</tr>
<tr>
<td>20/80 S/C</td>
<td>0.31</td>
<td>~60</td>
</tr>
<tr>
<td>0/100 S/C</td>
<td>0.27</td>
<td>~60</td>
</tr>
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</table>

Table 3. Changes in water/cement ratio, and initial- and final-setting times of 20/80 S/C ratio AACM and Class G well cement as a function of CMC content.

<table>
<thead>
<tr>
<th>CMC content, wt%</th>
<th>Water/cement, w/c, ratio</th>
<th>Initial setting time at 85°C, min</th>
<th>Final setting time at 85°C, min</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class G well cement</td>
<td>0</td>
<td>0.39</td>
<td>~90</td>
</tr>
<tr>
<td></td>
<td>0.4</td>
<td>0.48</td>
<td>~30</td>
</tr>
<tr>
<td></td>
<td>0.7</td>
<td>0.54</td>
<td>&gt;360</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>0.62</td>
<td>&gt;360</td>
</tr>
<tr>
<td>20/80 S/C ratio AACM</td>
<td>0</td>
<td>0.31</td>
<td>~60</td>
</tr>
<tr>
<td></td>
<td>0.4</td>
<td>0.38</td>
<td>~90</td>
</tr>
<tr>
<td></td>
<td>0.7</td>
<td>0.49</td>
<td>~120</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>0.52</td>
<td>~120</td>
</tr>
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</table>
Figure 1. Chemical structure of CMC.
Figure 2. Compressive strength of the specimens made with 40/60, 20/80, and 0/100 S/C ratios and Class G well cement after exposure in four different thermal environments.
Figure 3. XRD patterns for slag (a), Class C fly ash (b), 200°C-autoclaved 20/80 C/S ratio AACM (c), and autoclaved 20/80 S/C ratio AACM after heating at 300°C (d).
Figure 4. XRD pattern for autoclaved 0/100 S/C ratio AACM.
Figure 5. DSC curve of CMC.
Figure 6. FT-IR spectra for 80°C-made CMC film (a), and 200°C- (b), 250°C-(c), and 300°C-heated CMCs (d).
Figure 7. Changes in temperature of 20/80 S/C and Class G cement slurries after mixing with water at room temperature as a function of elapsed time.
Figure 8. Changes in compressive strength of 20/80 ratio AACM specimens exposed in different thermal environments as a function of CMC content.
Figure 9. Compressive strength vs. CMC content for Class G well cements after exposure in the different thermal environments.
Figure 10. Changes in porosity of 20/80 S/C ratio AACM exposed in the different thermal environments as a function of CMC content.
Figure 11. Porosity vs. CMC content for Class G well cements after exposure in the different thermal environments.
Figure 12. FT-IR spectra for slag and Class C fly ash as references.
Figure 13. FT-IR spectra for AACM with 1.0% CMC; 80°C-heated (a), 200°C-autoclaved (b), autoclave + 200°C-heated (c), autoclave + 250°C-heated (d), and autoclave + 300°C-heated (e).
Figure 14. Appearance of 200°C- heated 0 % (A), 0.4 % (B), 0.7 % (C), and 1.0 % (D) CMC-modified 20/80 S/C ratio AACMs after immersion in water.
Figure 15. In-situ exothermic temperature vs. elapsed time for 0.7 % CMC-modified AACM after beginning impregnation with water.
Figure 16. Changes in maximum exothermic temperature generated in AACM and Class G cement after impregnation with water as a function of CMC content, and visual comparison of magnitudes of self-degradation.
Thermal Decomposition of CMC

\[ \text{CMC} \rightarrow \text{CO}_2 + \text{CO} + \text{CH}_4 + \text{CH}_3\text{COOH} + \text{Na}_2\text{CO}_3 \]

Hydrolysis-Wet Carbonation in CMC-modified AACM

\[ (-\text{Na}_2\text{O}.\text{SiO}_2)_n^- + n\text{H}_2\text{O} \rightarrow n\text{Si(OH)}_4 + n\text{NaOH} \]

Non-reacted sodium silicate activator

\[ \text{NaOH} + \text{CO}_3^{2-} + 2\text{H}^+ \rightarrow \text{NaHCO}_3 + \text{H}_2\text{O} \]

Sodium bicarbonate

In-situ Exothermic Chemical Reaction

\[ \text{NaHCO}_3 + \text{CH}_3\text{COOH} \rightarrow \text{CH}_3\text{COONa} + \text{H}_2\text{O} + \text{CO}_2 \]

Sodium acetate

\[ \text{NaOH} + \text{CH}_3\text{COOH} \rightarrow \text{CH}_3\text{COONa} + \text{H}_2\text{O} \]

Figure 17. Self-degradation mechanism of heated CMC-modified AACM after contact with water.