Peculiar Set-Retarding Effect of Miserly Amounts of Pyrocatechol on Calcium Aluminate Cement Hydration

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Abstract

Pyrocatechol (benzene-1,2-diol) acts upon a calcium aluminate cement (CAC) as a very strong set and hardening retardant and a potent plasticizing agent at the same time. By studying CAC-pyrocatechol compositions using different analytical methods it was shown that pyrocatechol indeed slows CAC hydration and decreases strength of hardening stone. What makes this behavior a peculiar one is that pyrocatechol is known to accelerate Portland cement (PC) setting drastically.

Keywords

Pyrocatechol, Calcium Aluminate Cements, Hydration, NMR Spectroscopy

1. Introduction

Pyrocatechol (benzene-1,2-diol) has been reported to decrease Portland cement (PC) setting time significantly even when very small amounts of the substance was admixed [1]. In [2] an 27Al NMR spectroscopy disclosed this accelerating effect is due to a rapid ettringite formation by reaction between C₃A and gypsum, and that pyrocatechol has no noticeable effect on calcium silicates hydration (as shown by 29Si NMR spectroscopy). The primary factor that determines this acceleration effect is likely being an interaction between aluminum ions and pyrocatechol that results in chelate complex formation. Allowing for this theory, it’s intriguing to study how this particular phenol affects calcium aluminate cements (CAC) setting and hydration considering regular CAC primarily consists of CaO·Al₂O₃ and contains other calcium aluminates as well. When looked through dozens of publications this subject seems almost completely missed out.
The present work focuses mainly on pyrocatechol influence on CAC setting time, cement paste consistency and compressive strength of a cured cement paste, as well as how exactly pyrocatechol affects calcium aluminates hydration. We also attempt to describe and explain obtained results in correlation to known theories of CAC hydration.

2. Experimental Part

Aluminate cements used in this study are Secar 38R, Cement Fondu and Ternal EV (Kerneos, France). The XRD patterns of the dry cements are shown in Figure 1. The main phase of Secar 38R and Cement Fondu is monocalcium aluminate.
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(AC) with other phases also present, such as C_{12}A_{7}, C_4AF, C_2S, C_2AS. In case of Ternal EV (taken for comparison purposes), C_{12}A_{7} is a predominant phase; some amount of C_2A and C_4AF are present; the content of CA is inferior if at all.

Pyrocatechol was purified through evaporation and condensation (melting point 98˚C - 104˚C; 104˚C - lit.) and appears as colorless thin plates.

Water to cement ratio (w/c) in case of Secar 38R was 0.26, which refer to standard consistency of cement paste determined by GOST 310.3-76 (Vicat plunger penetrates to 5 - 7 mm from the bottom of the Vicat mold). In case of Cement Fondu and Ternal EV, the values of w/c were 0.265 and 0.21, referring to standard consistency of those cements, correspondingly. Pyrocatechol in amounts corresponding to (0.002 - 0.01)% by weight of CAC was dissolved in distilled water destined for mixing of cement.

To measure flow of cement paste a small cone of approx. 55 × 38 × 30 mm was used. The cone placed on a flow table was filled with cement paste (100 g), then the cone gets removed and the sample shaken 15 times.

Setting time of cement paste was determined using Vicat apparatus by GOST 310.3-76.

To evaluate the effect of pyrocatechol on cement paste hardening a compressive strength of 30 × 30 × 30 mm samples was measured. Samples were stored at 20˚C, 95% RH (normal conditions) for 24 hours, and then stored under water until the moment of measurement.

An early hydration of CAC Secar 38R with pyrocatechol admixture was studied by means of IR spectroscopy and ^{27}Al solid-state NMR spectroscopy. For this purposes a small amount of CAC paste with 0.01% of pyrocatechol was prepared by mixing the cement with water thoroughly. After 4 hours of storage under normal conditions a bit of sample (few grams) was powdered in mortar in an excess of acetone to remove any residue water and the sludge was vacuum filtered through a glass filter. The solid residue was then washed with acetone two more times and dried on glass filter. Dried sample had been stored in sealed container at −18˚C until the moment of measurements. A reference sample of pure hydrated CAC paste was prepared and handled the same way.

IR spectra were obtained using FTIR ATR spectroscopy (Shimadzu IRTtracer-100 FTIR Specphotometer with ATR accessory). High-resolution NMR spectra acquired on Bruker AVANCE II-500WB spectrometer. NMR spectra recorded at 130.32 MHz by single-pulse excitation, the pulse duration of 0.7 μs (π/12) with a delay of 0.5 sec, a number of scans was 2048. The samples’ material (approx. 100 mg) were packed in zirconia rotors (D 4 mm) and rotated at 13 kHz. Chemical shifts are given in ppm relative to AlCl₃∙6H₂O. An assignment of the signals is accomplished in accordance with published data [3] [4].

3. Results and Discussion

Physical and mechanical properties of Secar 38R cement paste with pyrocatechol admixture are shown in Table 1. It’s clearly seen that pyrocatechol slows CAC
setting dramatically with this effect more pronounced the more pyrocatechol is introduced. When introduced in larger amounts it also induces a strong plasticizing effect on cement paste and decreases strength of cement stone compared to reference samples. The retarding effect is observed also for Cement Fondu characterized by similar phase combination (Table 2).

In Figure 2 there are shown IR spectra of dry cement Secar 38R and cement pastes (both reference and with 0.01 wt% of pyrocatechol) at the point of 4 hours since water addition.

After 4 hours of hydration some noticeable changes take place in reference sample paste composition: an intensity of a signal at 800 cm\(^{-1}\), which refer to 4-coordinated aluminum ions of dry calcium aluminates [3] decreases, while signals of OH\(^{-}\) oscillation in hydration products at 1400 and 3400 cm\(^{-1}\) are quite distinct. At the same time, for pyrocatechol containing paste there’s no difference from dry cement spectrum which indicates that no hydration began yet.

This observation is also confirmed with \(^{27}\)Al NMR spectra (Figure 3): a strong peak at 80 ppm corresponds to AIO\(_4\) tetrahedra of aluminate phases of CAC (CA and C\(_{12}\)A\(_7\)) [4]. At 10 ppm there’s a weak signal that can be referred to aluminum-oxygen octahedra of C\(_4\)AF and C\(_4\)AS phases or it could also belong to hydrated cement as it absorbs air moisture.

| Table 1. Properties of cement pastes of Secar 38R with pyrocatechol. |
|------------------------|------------------------|
| **Property** | Pyrocatechol amount by weight of cement, % |
| Setting time, min: | 0.000 (reference) 0.002 0.005 0.010 |
| initial | 45 50 240 over 9 h |
| final | 165 260 580 - |
| Flow, mm | 55 56 100 100 |
| Compressive strength (MPa) at day: | |
| 1 | 54.3 54.5 41.7 1.7 |
| 3 | 65.0 - 51.6 - |
| 7 | 65.7 58.3 49.1 - |

| Table 2. Properties of cement pastes of other CACs with pyrocatechol. |
|------------------------|------------------------|
| **Property** | Cement Fondu  Ternal EV |
| Pyrocatechol amount by weight of cement, % | 0.000 (reference) 0.005 0.010 0.000 (reference) 0.005 0.010 |
| Setting time, min: | |
| initial | 85 85 over 8 h 40 40 60 |
| final | 240 over 8 h over 20 h 120 70 100 |
| Flow, mm | 64 64 123 182 195 202 |
At 4 hours from the moment of water addition a reference paste has fully set, meanwhile for pyrocatechol containing sample it hasn’t even started. Strong differences between spectra of those two can be observed: for reference paste calcium aluminate phases’ signal diminishes and a strong peak of hydrates arises at
10 ppm, when pyrocatechol stops cement hydration almost completely. Comparing integral intensities of signals at 80 ppm allows for approximate quantitative estimation of hydration rate for both cases, which results in 23% for reference and about 1% of pyrocatechol sample.

Summarizing results of mechanical properties evaluation and spectroscopic analyses it may be concluded the pyrocatechol suppresses hydration of CAC, its setting and hardening and while this effect is not something completely surprising (considering many organic compounds, in particular α-hydroxy carboxylic acids, carbohydrates and so on behave the same way) there are two features that takes pyrocatechol aside. First, it influences Portland cement and calcium aluminate cement in very different ways: accelerating the setting of the former [2] and slowing for the latter. Secondly, these effects become noticeable at miniscule dosages of pyrocatechol—some hundredths and thousandths of a percent by weight of cement. Widely known and used in practice organic modifiers of setting time usually affects both PC and CAC in the same manner and their dosages typically lay in range of tenths of a percent (see, for example, [5]).

To qualitatively study how various components of PC reacts with water in the presence of pyrocatechol, additional experiments was carried out using several materials: an industrial grade Portland cement clinker (phase composition, wt%: C₃S 54-57, C₂S 19-21, C₃A + C₄AF 24-25 – petrographic examination), synthetic C₃A and C₄AF minerals and a calcined gypsum (β-calcium sulfate hemihydrate). These materials in finely powdered form were mixed with distilled water (as reference samples) at a ratio of 0.4 (0.5 for CaSO₄·0.5H₂O) and with water solution of pyrocatechol (0.02 wt% by the weight of dry powder) then reaction results were examined organoleptically. Several observations were made: in comparison to reference samples pyrocatechol 1) exerts a strong plasticizing effect upon gypsum paste, 2) leads to a very strong exothermic effect and an instantaneous set of C₃A, 3) has no influence on C₄AF hydration, 4) accelerates PC clinker set and heat release as well.

All the above mentioned allows for the conclusion that the quickened set of PC with pyrocatechol admixed is mainly due to its influence on C₃A phase hydration kinetic, which agrees well with our previous research results [2]. Hence the pyrocatechol is expected to have an essentially different impact on hydration of tricalcium aluminate of PC and those aluminates of CAC, such as CA, which are of less basicity.

To give a comprehensive explanation of this phenomenon is quite difficult considering an actual mechanism of dissolving of calcium aluminates isn’t worked out yet and it’s still debated be it congruent or incongruent [5] [6] [7] [8]. Based on available data on early period of hydration of different aluminate phases we can only make some assumptions about those effects.

When first come into contact with water, C₃A particle surface is overgrown with hydration products which limit its further reactivity (both with presence of gypsum or without it) for at least some period of time [7] [8]; for the duration of the period a high hydraulic activity of C₃A is extinguished. This thin layer of
products is proposed to consist of unstable hexagonal phases, which then
verse into stable state. This process increases permeability of the layer so the 
hydration goes on and heat is been released [6] [8]. Hence such compounds that
stabilize initial layer slows the hydration and vice versa [6] [7], e.g. triethanolamine
(TEA) is known to promote C₃A reaction with water [8].

Decrease in hydration rate of C₃A can also be induced by formation of
amorphous hydrated aluminum hydroxide layer covering the grain (perhaps, 
due to incongruent dissolving of C₃A). Its potency to screen the grain of C₃A
from water depends on many factors, including composition of liquid phase [5]
[6] [7]. Followed by layer destruction in which Ca²⁺ and OH⁻ ions from liquid
phase also participate, C₃A hydration eventually resumes.

In presence of gypsum some amorphous layer of Ca²⁺ and SO₄²⁻ stabilized
product is formed on surface first, which slows hydration and initiate the induc-
tion period, and only then from this primal product ettringite crystals start to
grow [7]. This way a typical PC hydration rate and setting rate are controlled
predominantly by Ca(OH)₂ and C-S-H nuclei formation [5]. Consequently, as
Ca²⁺ and SO₄²⁻ in liquid phase exhaust, the layer becomes unstable and permea-
able so C₃A surface is not restricted from water access anymore.

On the other hand, if excess ettringite formation is promoted somehow in
“C₃A + gypsum” system it may shorter the induction period and expedite ceme-
tment setting. In practical applications this effect is usually achieved by intro-
ducing either some aluminum containing compounds that readily react with
gypsum themselves to form an ettringite [9] or other additives to induce the
reaction between PC own phases and gypsum—one example of the latter is py-
rocatechol [2].

Pyrocatechol is a chelating agent that binds calcium and aluminum ions into
strong water-soluble complexes. This interaction seems to affects primal layer of
hydrates that inhibit C₃A or perhaps the layer formation is cancelled in pyrocate-
thol presence, so C₃A is unleashed to react with water whether gypsum is
available or not. TEA is known to possess similar properties, so its action can be
explained using similar approach.

Calcium aluminate (CA)—a main constituent of CACs with moderate Al₂O₃
content—reacts with water via through-solution mechanism [7], first proposed
by Le Chatelier and confirmed by numerous researches thereafter [5], although
there still appears a layer of aluminum-rich hydrated product when CA phase
makes contact with water [7]. CA likely undergoes congruent dissolution, this
leads to supersaturation of solution by AH₃ and hydrated calcium aluminates
and create favorable conditions for crystallization nuclei to emerge and grow. In
this case, the duration of induction period is determined by the time it takes
these nuclei to become stable and start growing. By the end of induction period
setting occurs. C₁₂A₇ phase, if presented, dissolves readily and quickly supersa-
turate the solution with respect to C₃AH₆ so nucleation rate increases considera-
tably [7].

In those cements where CA predominate, pyrocatechol may act as “poison-
ing" agent to inhibit nuclei formation by capturing calcium and aluminum ions from solution into strong complexes. In this regard, pyrocatechol behave as a typical retardant, however the retarding effect is pronounced even when miniscule amounts of pyrocatechol are introduced, and what we call a miniscule is actually an order or two below dosages of widespread retardant agents used in practice nowadays.

In view of all this, a specific note should be made about Ternal EV behavior. Composed principally of С12А7, that “stands between” C_3A and CA on CaO-Al_2O_3 phase diagram, and some amount of C_3A (as is identified by XRD) and traces of CA (if any), this cement shows no clear behavior when exposed to pyrocatechol, at least for dosages tried in this work (Table 2).

4. Conclusions

As it was disclosed by spectral analysis and mechanical properties evaluation tests, the pyrocatechol inhibits hydration of calcium aluminates of CA, which in turn affects setting time of cement paste and its compressive strength development. At the same time, pyrocatechol accelerates Portland cement hydration by promoting C_3A reaction with water and gypsum and inducing ettringite formation (C_3A + 3CaSO_4 \cdot 2H_2O + 26H_2O \rightarrow C_3A \cdot 3CaSO_4 \cdot 32H_2O).

Set-retarding effect of pyrocatechol is exhibited at miserly dosages, some 0.002 - 0.005% proportional to calcium aluminate cement weight. The pyrocatechol binds calcium and aluminum ions into strong chelate complexes and thus inhibits nuclei formation and developing.

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