The Influence of Citric Acid on Setting Time and Temperature Behavior of Calcium Sulfoaluminate-Belite Cement

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A. INTRODUCTION

Concrete is the most widely used manmade material worldwide for reasons that range from cost effectiveness to easiness to mold [1]. Nonetheless, research is still needed in order to improve properties such as strength, durability, chemical resistance, and to reduce its environmental impact. Many alternative cements including calcium sulfoaluminate (CSA) cement have been developed. A major advantage of some alternative cements is a reduction in the amount of carbon dioxide released during cement manufacture. CSA cement production releases about half the amount of carbon dioxide (49% less) that ordinary portland cement releases [2]. Other advantages of CSA cement include rapid setting time, higher strength than portland cement over time, rapid strength gain, and improved durability [3]. This paper focuses on one of the special types of the CSA family, CSA-Belite, and its rapid setting property. This type of cement has additional properties such as near-zero shrinkage, immunity to sulfate attack, and lower porosity, among others [12]. According to Dr. E. Bescher from the Department of Materials Science and Engineering at University of California Los Angeles, the unit price ratio for BCSA to portland cement is close to 1.4 [12]. Other sources claim that CSA cement is about four times more expensive than portland cement type-I [3]. However, despite being more expensive, BCSA can have a longer life than portland cement, resulting in a 53.5% reduction in cost. This cost saving is based on an airport paving project at Seattle Tacoma Airport [12]. BCSA cement has the ability to harden at an accelerated rate, which is beneficial for construction time frames if managed correctly. Consequently, the rapid setting time (20 minutes) of BCSA cement is a challenge that has limited its application in the real world. One of the most common solutions has been the implementation of citric acid as a retarder [2]. This research investigated the effects of citric acid on the setting time of BCSA cement with the purpose of finding solutions to constructability problems. Furthermore, this research will also present and study the collateral effects on the compressive strength and the temperature behavior with the help of not only empirical data but also with a theoretical perspective.

B. THEORY

CSA-Belite Chemistry

The chemical composition of BCSA cement differs considerably from portland cement and this can be observed in its physical behavior such as setting time, color, heat generation, etc. Portland cement is composed of three main chemical compounds: silica (SiO$_2$), calcium oxide or lime (CaO)
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and alumina (Al₂O₃) [10]. In general, these compounds are present in the form of other configurations as shown in Table 1. This table also shows the main products of the cement-water reaction. Other products such as calcium silicate hydrate (C-S-H) appear in the structure at later ages [18]. The following table presents the chemical formulae of the components of CSA, BCSA and type-I portland cement [3],[4],[12]. Type- I portland cement has been included in the table since it is the most widely used type of cement [10].

<table>
<thead>
<tr>
<th>Reactants</th>
<th>Compound</th>
<th>Cement Chemist Notation</th>
<th>Percentage by mass (%)</th>
<th>Type I</th>
<th>CSA</th>
<th>CSA-B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ye’elimite</td>
<td>(CaO)₄(Al₂O₃)₃SO₄</td>
<td>C₄A₃S</td>
<td>---</td>
<td>50-60</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>Alite</td>
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<td>C₃S</td>
<td>59</td>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td>Belite</td>
<td>(CaO)₃SiO₂</td>
<td>C₂S</td>
<td>15</td>
<td>0-20</td>
<td>45</td>
<td></td>
</tr>
<tr>
<td>Ferrite</td>
<td>(CaO)₄(Al₂O₃)Fe₂O₃</td>
<td>C₄AF</td>
<td>8</td>
<td>5-15</td>
<td>2</td>
<td></td>
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<tr>
<td>Aluminate</td>
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<td>C₃A</td>
<td>12</td>
<td>---</td>
<td>---</td>
<td></td>
</tr>
<tr>
<td>Anhydrite</td>
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<td>CS</td>
<td>2.9</td>
<td>15-25</td>
<td>15</td>
<td></td>
</tr>
</tbody>
</table>

**Main Products**

<table>
<thead>
<tr>
<th>Ettringite</th>
<th>(CaO)₆Al₂O₃(SO₄)₃(H₂O)₃₂</th>
<th>C₆AS₃H₃₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monosulfate</td>
<td>(CaO)₄Al₂O₃SO₄(H₂O)₁₂</td>
<td>C₄ASH₁₂</td>
</tr>
<tr>
<td>Calcium Aluminate</td>
<td>Al(OH)₃</td>
<td>Al(OH)₃</td>
</tr>
</tbody>
</table>

Table 1. Chemical composition of Portland, CSA and Belite-CSA cements [3],[4],[12].

Each one of these compounds contribute in a different way to the hardened properties of concrete. C₃S and C₂S are responsible for the hydration rate and gain in strength of portland cement in the early and late stages respectively. These compounds cause concrete strength gain primarily through the formation of C-S-H. For CSA, C₄A₃S is the responsible for the rate of hydration, expansion, and rapid grain strength of concrete due to the amount of alumina (Al₂O₃) it contains [10]. Table 1 shows that BCSA has a much greater amount of alumina than portland cement, which results in faster strength gain and refractory properties. Another important compound is C₃A, which contributes in the increasing of heat of hydration, but reduces its resistance to sulfate attack and volumetric shrinkage [10]. In BCSA, rapid formation of ettringite is responsible for the rapid strength gain and short setting time [3].
The chemical reaction of the cement paste/concrete can be represented by five equations according to Burris and Kurtis [4]. These equations are helpful to demystify the overall process because having multiple reactions means different reactants are consumed at different rates. Since there are multiple reactions, multiple reaction rates can be studied, but this research focuses on studying the influence of citric acid on the overall reaction rate. The assumptions made for the overall process can also be applied to each one of the five chemical equations as demonstrated in the following subsection.

Reaction Kinetics – Maturity Method

The reaction between cement and water can be described as heterogeneous since there is a phase change. The rate of reaction for this type of reaction is affected by parameters such as temperature, pressure, composition, mass transfer, and heat transfer [6]. Hydration rate is the reaction rate that yields the binding properties of hardened concrete after mixing its components [10]. This cement-water reaction goes infinitely as long as enough water is provided [10]. Chemically, this rate can be defined as the rate of moles formed of a specific product (ettringite for the case of BCSA cement) over the volume of fluid, and time [6]. Measuring the reaction rate based on this definition is not possible unless more specialized techniques such as mass spectrometry and quantitative phase microscopy are used. Nonetheless, it is possible to find a correlation between the citric dosage used and the rate of reaction through the maturity method and Arrhenius equation. The temperature-time plots can be used to compare the results of the rate of the reaction to setting time. The period it takes for the cement paste to go from fluid to completely solid is an arbitrary metric generally defined as setting time. Since the reaction rate and setting time are inversely proportional, both should yield linear trends [7]. The kinetics of any chemical reaction can be described with the Arrhenius equation and the rate of chemical reaction equation [6],[7].

\[
    k_t = Ae^{-\frac{E}{RT}} \quad \text{Arrhenius Equation, Equation 1}
\]

\[
    v = k_t [A]^n [B]^m \quad \text{Rate of Reaction, Equation 2}
\]

In Arrhenius equation, \(k_t\) represents the rate constant, \(A\) is a frequency constant, \(R\) is the universal gas constant (8.314 J/K-mol), \(T\) is the measured temperature, and \(E\) is the activation energy. In the second equation \([A]\) and \([B]\) represent the concentration of reactants and \(n\) and \(m\) are the reaction orders. These concentrations can be considered as constant for this research since the
water-cement ratio remained the same through the experiment. The Arrhenius equation can be used to establish the relationship between the setting time and rate constant if the ASTM C1074-17 standard is used. However, an alternative method which relates setting time to age development will be used. This method is described by Hover et al, and it defines equivalent age as the time it takes for the specimen to reach a certain level of strength development based on the temperature-time behavior. The maturity function by Freisleben-Hasen and Pederson is derived from the Arrhenius equation [7].

\[ t_e = \sum_0^t e^{-\frac{E_a}{R} \left( \frac{1}{T_a} - \frac{1}{T_{ref}} \right) \Delta t} \]  

Maturity function, Equation 3

In the maturity equation, \( T_a \) is the average temperature over time, \( T_{ref} \) is the reference temperature, and \( t_e \) is the equivalent age at reference temperature. For this case, the development age of each specimen can be obtained with its corresponding final setting time and temperature. The slope of \( \ln(k_t) \) vs \( 1/T \) is a factor of the activation energy. The variable \( k_t \) is replaced by the inverse of setting time since it is unknown and it is proportional to setting time.

Comparison of Vicat and Penetration Resistance Test

The Vicat and Penetration resistance test are methodologies utilized to measure the initial and final setting time of mortar or cement paste. Initial set time can be defined as the period it takes to stiffen (or lose its plasticity). Final time is defined as the time it takes the mortar or cement paste to harden, or reach structural strengths [10]. The Vicat apparatus measures these times as a function of the displacement of a needle into the mortar. The penetrometer measures these times as a function of the stress created after pushing a series of needles over a sample. Given that the 1 mm-needle pushes a weight of 300 g into the paste, the resulting stress is 536 psi (3.7 MPa), which is close to the stress indicating initial set in the penetration resistance test, 500 psi (3.4 MPa). This means that both tests should have an acceptable level of agreement in their initial set times [9].
For the analysis of the temperature behavior, it is necessary to establish a relationship between the initial and final setting time and the temperature at these points. For this analysis, the setting time obtained with the penetration resistance test will be used because the temperature was measured in the sample where the penetration resistance test was performed. In addition, there are some reasons to believe that the measurements provided by the penetration test are more reliable. First, the Vicat test does not provide a strong relationship between stiffening and setting time. In a study performed by Lee and Hover [8], the Vicat test did not provide any measurement of stiffening before the initial set time of the same paste compared to the penetration resistance test. This study also concluded that the penetration resistance test is the most accepted method, which provides an accurate correlation between setting time and the stiffening of the paste [8]. Both tests are based on the same physical principle of deformation over a surface area. A study of the Vicat apparatus led by Sleiman et al. showed that different needle masses yielded different results. They concluded
that the assumption (quasi-static penetration) on which the Vicat test relies are not suitable during the thixotropic period, which is approximately the first hour [16]. A quasi-static process is a process that occurs slowly. Another important aspect to consider is that for the penetration resistance test, the mass of the batch was greater, and consequently more heat is produced, which speeds up the reaction. This is more realistic to what happens in the field, and one of the reasons why the setting time was shorter for the penetration resistance test almost all the time. Despite the Vicat test’s deficiencies, it is an acceptable technique since the needle shape and frustum depth produce normalized results that can be used to compare setting times between cements prepared in the same fashion [16]. Consequently, this research will also consider the results from the Vicat test since it is widely utilized by cement manufacturers to measure setting times [17].

Effect of Citric Acid on Dynamic Viscosity
The influence of citric acid on the cement paste extends to other properties, not just setting time. It improved its fluidity, which can be represented through dynamic viscosity. Viscosity can be defined as the flow resistance of a fluid. The viscosity of a citric acid aqueous solution can be calculated with one form of the Arrhenius equation [19].

\[ \mu = \mu_0 e^{-\frac{E_a}{RT}} \]  

\( \mu \) represents the dynamic viscosity of the solution, \( \mu_0 \) is the dynamic viscosity of water, \( R \) the gas constant, \( T \) the absolute temperature, and \( E_a \) is the activation energy. For this research, the viscosity of water and temperature will be treated as constant since the variation of these parameters is small. The only variable is activation energy, which increases as more citric acid is used in the mix. Negative catalysts or retarders increase the activation energy, which reduces the reaction rate [20]. Therefore, increasing the activation energy, reduces the viscosity of the mixture according to equation 3. If the viscosity of the mixture is reduced, the flow resistance is reduced as well, which results in an improved and more fluid mix. The viscosity of the mixture was influenced by the dosage of citric acid and not by the water-cement ratio since it was kept constant. Equation 4 shows that the only factor of water that affects the viscosity of the mixture is the viscosity of the water.
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C. EXPERIMENTAL PROCEDURE

There are three parameters of BCSA cement being studied in this research: setting time, compressive strength, and temperature behavior. Therefore, the procedures to measure each one of these will be described separately. These procedures follow the methods in accordance to the ASTM standards with some minor modification due to the nature of BCSA cement.

Cement Paste

The cement paste was made in accordance to ASTM C305-14 [13]. First, the water (mixed with the citric acid retarder) was placed in the mixing bowl before adding the cement so that the citric acid was well dispersed in the water. The water to cement ratio (w/c) in this experiment was 0.48. Other specialized studies related to the effect of retarders on CSA cements used w/c values of 0.4 or 0.5 [4],[5]. CSA cements generally requires a higher w/c than portland cement in order to fully hydrate [3]. The dosage of citric acid measured by number of fluid ounces per 100 lb. cement was varied: 0, 4.5, 9, 13.5, 18 and 36 (added later in the research). In order to obtain these dosages, 5 lb. powdered citric acid was added to 1 gallon of water to make a solution similar to a typical concrete admixture. Based on these ratios, the amounts of citric acid were: 0.0%, 0.18%, 0.35%, 0.53%, 0.70% and 1.4% by weight of cement. After adding the cement, the components were left to absorb for 30 seconds. Then, the laboratory bench mixer was started at low speed for a time between 30 seconds and 1 minute. After this time, the mixer was paused for about 15 seconds to scrape the bottom of the mixing bowl to make sure all the components were being mixed appropriately. Then, the medium speed was set for a time between one to two minutes. This is a slight deviation from the ASTM C305-14, which is based on a mix of 650 g of cement with a water to cement ratio determined by ASTM C187-16. These deviations are justified with the fact that the amount of cement utilized was more (6166.65 g) and the w/c ratio was established to be 0.48 since it is more common for this type of cement [4]. The amount of cement paste made by batch was approximately 308 in³ (5047.2 cm³). The mixing water was kept at a temperature within a range of 73.5 ± 3.5 °F (23.0±2.0 ºC). Tests were performed at 21.1º, 23.3º and 25º Celsius specifically to see the variability of the results based upon water temperature.

Setting Time

The initial and final setting time were determined by means of two methods. The setting times were determined in accordance to ASTM C403/C403M-16 [15] and ASTM C191-18 [14]. Both
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methods were performed inside an environmental chamber where the ambient temperature was kept between 73.5 ± 5.5 °F (23.0 ± 3.0º C). Due to high temperatures during the summer, the ambient temperature in the environmental chamber was higher than the temperature of the range sometimes. Also, the humidity was kept above 50% as the ASTM standards dictates. It was kept between 65% and 75%, specifically. The procedure for each methodology will be explained as follows.

I. Vicat Needle (ASTM C191-18)

Once the cement paste was made, part of it was placed in a non-absorbent conical ring with an internal volume of approximately 14 in³ (229.4 cm³). The conical ring was placed over the glass plate, and the cement paste was poured through the smaller diameter of the ring until the paste surface and the ring border were leveled. The surface was leveled and smoothed carefully with the help of a rod to avoid adding stress. For the specimen without retarder, the ring was tapped on the sides to make sure voids are filled with the paste. This was not necessary when the retarder was used because the mix was more fluid. The mix was more fluid despite keeping the water-cement reaction constant for reasons explained in the literature review. Immediately after molding, the specimen was placed in the environmental chamber for testing. This test was performed with a manual Vicat apparatus, which is described as Method A in the ASTM standards. The indicator was placed at zero before releasing the rod and the one-millimeter needle into the paste. The first penetration of the needle into the paste was done at the center of the specimen, and the subsequent measurements were done approximately five millimeters away from the previous penetration. The pattern of these measurements was circular and moving out of the center. Measurements were performed at least 10 mm away from the border.

Depending on the nature of the specimen, the time between the preparation of the specimen and first measurement varied. Likewise, the measurements were done at different time intervals depending on the amount of citric acid used. For example, if no retarder was used, faster measurements were required to achieve an accurate view of the setting behavior. On the other hand, if more citric acid was used the measurements were spread out to capture the range of behavior between mixing and setting. Data points were collected until zero displacements of the needle or no marks were observed on the cement paste within a period of 90 seconds. The time at which this happened is the final setting time according to ASTM
C191-18. The initial time was determined by linear interpolation of the time at which 25 mm of displacement is believed to occur. The data of this test were plotted in a set of piecewise linear functions where the function of interest was the one crossing the 25 mm displacement.

II. Penetration Resistance (C403/C403M-16)

In contrast to the Vicat test which has a criterion based on the displacement of the needle, the Penetration Resistance test has a criterion based on the stress created when some force is applied. First, approximately 170 in$^3$ (2785.8 cm$^3$) of cement paste was poured into a cylinder leaving a gap of one inch from the cylinder’s top edge. The cylinder had a diameter and height of 6 in. and was made of a non-absorbent plastic material. The surface of the sample was smoothed with a rod. The same rod was utilized to tap the sides of the container to improve consolidation. Then, the specimen was placed in the environmental chamber for testing. This test consists of pushing six needles of decreasing diameters vertically one inch into the specimen, resulting in six different stresses. The measurements were performed at least 1/2 in. apart, and 1 in. away from the border. Due to limitations of the specimen surface and the need to perform additional penetrations these separation distances were less sometimes. However, if that was the case and the data point was misrepresentative, it was discarded for calculations purposes.

At least six measurements were performed, conforming to ASTM C403. According to the standard, the initial setting time occurs when the cement paste has reached 500 psi (3.4 MPa) in strength, and the final setting time occurs when the penetration strength is 4000 psi (27.6 MPa). The data is presented in two different ways. First, a simple plot of time versus stress will yield a curved line. Nevertheless, straight lines help to have a better interpretation of the nature of the experiment. Therefore, the standards give guidance on how to present the data points in a log-log plot. Despite the two ways to present the data, the results for setting times should be close in value. It is important to highlight that the required ambient temperature varies slightly compared to the one required for the Vicat test. Nonetheless, for consistency the ambient temperature was kept at 73.5 ± 5.5 °F (23.0 ± 3.0° C) as previously mentioned.

Compressive Strength and Temperature Behavior

The rest of the cement paste was used to mold 12 2 in. cubes, which were tested ten minutes after both setting time tests were done then at three hours, one day and seven days. Due to the extremely fast setting (< 20 minutes) of the specimen when no retarder was used, the setting time samples
and the cubes were cast from different batches. This only applied for the specimens with zero citric acid and water temperatures of 23.3°C and 25°C. The cubes were tested in a compression machine according to ASTM C109 [22].

The temperature change within the cement paste was measured by inserting a type-K thermocouple into the center of the penetration resistance specimen contained in the 6 in. cylinder. This thermocouple was coated with epoxy to avoid issues if contacted with water. The temperature was recorded with an HBM QuantumX data acquisition system at a frequency of 10 Hertz.

**D. RESULTS AND ANALYSIS**

**Setting Time**

The results from the Vicat test and Penetration resistance test yielded linear trends. This means that as more citric acid dosage is used, the increase in setting time behaves linearly. The data points at the dosage of 36 oz/100 lb of cement were not considered in this trend because the ambient temperature at which these measurements were performed was lower compared to the first five dosages. The R-square values of these plots are close to one, which strongly suggest linear relationship between setting time and dosage of citric acid.

![Figure 2. Setting Time dependence on dosage of citric acid (Vicat Test).](image-url)
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The tests performed have different mechanisms and levels of reliability as previously discussed. However, they both showed that the relationship between setting time and dosage of citric acid is linear. Moreover, if the slopes of the initial and final setting time trends of the Vicat test results are compared to their equivalent on the penetrometer test results, it shows that their magnitudes are close. For both setting time tests, the difference between first and final setting time increases as the dosage of citric acid increases. For the Vicat test, the differences in time between the initial and setting time are 2.1, 5.4, 14.8, 17.8 and 23.8 minutes for its corresponding dosage. Therefore, the linear trend lines diverge as the dosage increases. This divergence is also observed in the penetrometer test. This divergence is corroborated with the difference of slope or by analyzing the separation between initial and final times as the dosage increases. The ratio between consecutive initial and final differences for consecutives dosages can describe the level of divergence. This ratio starts at 2.6 and finishes at 1.3 for the Vicat Test. For the Penetrometer Test, this ratio starts at 2 and finishes at 1.4. If this ratio is one, it means that the difference between initial and final time is constant, and if it is below one, this gap is being reduced and the linear equations converge. The results show that these ratios are always above one, meaning that the gap always increases, but at some point it approaches one. The linear trend lines of these ratios can be utilized to extrapolate the setting times for higher dosages. An extra dosage of 1.4% of citric acid by weight

Figure 3. Setting Time dependence on dosage of citric acid (Penetrometer Test).
of cement was tested four months after (December). The results of the 36 oz/100 lb of cement (1.4%) are shown in the graphs, but they are not considered in these calculations because of the different ambient temperatures and humidity at which these specimens were tested. Even though these specimens were tested at temperatures that fall within the allowed range of the ASTM, they were not included to keep consistent data. Furthermore, setting times of 100 minutes were measured with 18 fl. oz. of the citric acid solution. These times are more consistent with the setting time of portland cement, but for most applications higher setting times than this are not needed.

**Maturity Method Correlation to Setting Time**

Calculations using the maturity approach were performed and results are represented in the following Figure 4.

![Figure 4. Equivalent age vs dosage of citric acid.](image)

The results from Equation 3 in the maturity method yielded a linear relationship between equivalent age and citric acid dosage as well. This strengthens the hypothesis of linearity between citric acid dosage and setting time. The data used for these calculation are in Table 2. Also, \( T_{ref} \) was set as 20°C (293.15 K) as recommended by the ASTM C1074-17 [21]. \( T_a \) represents the average temperature of the measurements until final setting time was reached. \( T_s \) represents the temperature at final setting in the penetration resistance test. \( E_a \) is a factor of activation energy calculated with the penetration resistance final setting times as explained in the theory section.
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<table>
<thead>
<tr>
<th>Dosage</th>
<th>Initial Set ( t_i ) (min)</th>
<th>Final Set ( t_f ) (min)</th>
<th>( t ) (min)</th>
<th>( \ln(1/t) )</th>
<th>( T_a ) (K)</th>
<th>( 1/T_s ) (1/K)</th>
<th>( E_a ) (J/mol)</th>
<th>( t_e ) (+)</th>
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</table>

Table 2. Data and results from the maturity method to obtain equivalent age, \( t_e \).

There are some deviations with respect to Hover’s work. In this research, the equivalent time is obtained as a function of curing temperature and citric acid dosage. The temperature values used correspond to the temperature values at its respective setting. Moreover, a coefficient factor is required to obtain the actual activation energy and equivalent time, but the obtained values can be used to compare behaviors for setting time and equivalent age as a function of citric acid dosage. These values of equivalent age help to confirm that the setting time is linearly dependent on the citric acid dosage. Figure 4 shows that the time it takes for a specific specimen to develop final setting time conditions increases linearly as citric acid is added. There is some level of uncertainty due to the limited amount of data points available, but it generally confirms the trend observed in the Vicat and penetration tests, even including the 36 fl. oz. citric acid dosage, which was performed at a different ambient temperature.

Compressive Strength

The results for compressive strength are analyzed in accordance to two parameters. First, the dosage dependence because it is the aim of this research and second, the water temperature dependence. Figure 5 shows that adding citric acid does not compromise the compressive strength of BCSA cement mortar at after setting and one-day stages except for the 36-oz. dosage. Adding
citric acid to the mix increases the compressive strength slightly at the seven-days stage as it can be observed for the 36-oz. dosage. Furthermore, Figure 6 shows that there is not a significant impact of the temperature ranges of the mixing water used in this study on the compressive strength. Nonetheless, hotter water increases the rate of reaction and reduces the setting time.

![Figure 5. Compressive strength results with its corresponding citric acid dosage.](image)

![Figure 6. Compressive strength dependence on water temperature.](image)
Specimens prepared with portland cement type I can exhibit a minimum of 1800 psi (12.4 MPa) and 2800 psi (19.3 MPa) strength at 3 and 7 days respectively [10]. The specimens made of BCSA cement presented an average strength of 3557.5 psi (24.5 MPa) and 4967.0 psi (34.2 MPa) for 3 and 7 days. Moreover, concrete made of BCSA cement is expected to have a higher strength since the strength of concrete depends on the strength of the aggregates and the cement paste.

Fractures were noticed in the compressive strength cubes when no citric acid was used. These fractures can be attributed to the high temperature gradient. This is explained with more details in the temperature behavior section. The cubes tested for 1-day and 7-day compressive strength presented cracks even though they were cured by two different methods. First, they were placed in a water tank 60-90 minutes after the final setting time for curing, and they presented cracks at the moment they were tested. These specimens could not be tested for compressive strength. Therefore, the mortar specimens were cured in air alternatively, and even though they still presented smaller fractures, they were able to hold load, so they were tested. The other specimens containing citric acid did not present fractures. This may be a limitation of testing a pure cement paste, since the heat generation is very high when no aggregate is used and no citric acid is present.

**Temperature Behavior**

This section focuses on how the increase in citric acid dosage affects the temperature behavior of the cement paste during and after setting. The temperature behavior will be studied by identifying parameters such as point of maximum temperature, point of maximum change of temperature and the area under the curve of the temperature-time plot (related to the heat transfer) for each one of the different dosages. To measure heat more accurately, other parameters such as mass transfer and heat capacity need to be considered. Additionally, this section will study the special case of no citric acid since cracks were observed after curing. The following plots represent the measurements of the temperature versus time during the cement reaction up to a few hours after setting time. Figure 6 shows the temperature-time plots for the varying dosages of citric acid set retarder.
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Figure 6 shows that the peak temperature decreases and is shifted in the time axis as more citric acid is added. The points in time at which the maximum temperature occurred were calculated by averaging the last and first point at which a maximum temperature was detected since multiple maximum values were measured due to the frequency of the apparatus (10 Hz). Furthermore, the points in time at which the maximum slope occurred for each of the temperature-time plots were calculated using a MATLAB function. These points are represented with black dots in Figure 7. The results for maximum slope yielded a linear trend with the equation $y = 4.6127x$ and an R-square value of 0.9671. The maximum change (slope) in temperature is represented by $y$ and $x$ is the dosage of citric acid.
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Figure 8. Initial setting time and maximum slope time as a function of citric acid dosage.

Figure 9. Average temperature, maximum temperature and time of maximum temperature results.

It can be observed from Figure 9 that the maximum temperature is reduced by a linear factor as citric acid is added. Despite this fact, the time at which the maximum temperature occurs increases
because the reaction has been decelerated. All the temperature parameters studied in this section have yielded linear trends.

These temperature plots can be utilized to explain the reasoning behind the slight increase of the compressive strength due to increase in citric acid dosage. First, this correlation can be explained by means of the maturity method. This concept explains that early strength is higher when temperatures are high during the curing process, and late strength is higher when temperatures are low during the curing process. The slow hydration rate and low temperature allows the cement paste to maximize the reduction of porosity due to a uniform formation of ettringite and other reaction products [1]. Besides, the slow hydration, fluidity of the cement paste could have also been a factor in the development of strength. The mixing proportions and properties of the materials used for the cement paste influence the consistency of the mortar [10]. In this case, as more citric acid was added to the mix, the consistency of the mix was improved. Consequently, the material was easier to mold, fill space and reduce the porosity. The compressive strength increases as the percentage of air voids in concrete is reduced [10].

The results of the compressive strengths somewhat match the maturity theory. The dosage at which the early rapid gain strength is reduced lies between 0.7% and 1.4% of citric acid because the 1.4% citric acid specimens showed lower strength at the after setting and three hours’ stages, and higher strength at the seven-days stage compared to the 0.7% specimens. Despite the fact that this theory cannot explain the small variations of compressive strength for specimens with dosages below 0.7%, the idea of this theory helps to understand that at some dosage of citric acid the early rapid gain strength property is reduced since the temperature and dosage are inversely related. This deficiency can be attributed to the fact that the maturity approach does not consider other factors such as early age curing temperatures and humidity [1]. The maturity approach also establishes that time and temperature of hydration have a correlation with strength. This is expressed as a maturity function which is the area under the curve of the temperature-time plot. The results of the area under the curve are presented in the following table:

<table>
<thead>
<tr>
<th>Dosage (%)</th>
<th>0.0</th>
<th>0.18</th>
<th>0.35</th>
<th>0.53</th>
<th>0.7</th>
<th>1.4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Area under the temperature-time plot (°C·min) (10^3)</td>
<td>1.5717</td>
<td>2.2064</td>
<td>2.5308</td>
<td>3.0628</td>
<td>4.0403</td>
<td>5.2457</td>
</tr>
</tbody>
</table>

Table 3. Area under the curve of temperature-time plots.
The table shows that the area under the curve increases as the dosage of citric acid increases. The correlation is that as the area increases, the compressive strength should increase [1]. However, the results of this research show that this fact only applied to the strength at 1 day or more since the 1.4% dosage specimens had lower compressive strengths after setting and at 6 hours despite having greater area under the temperature-time plot. Further studies are required to understand the mechanism behind the formation of ettringite at early stages when high dosages of citric acid are used.

One of the physical results on the cement paste was the appearance of cracks developed during curing. The cubes of cement paste were placed in a water tank approximately one hour after it reached the peak temperature. Fractures only appeared on the cubes that did not have citric acid. Smaller cracks were also observed in samples which were cured in air. This cracking might be attributed to the evaporation of water at high temperatures. The mortar cubes developed strength while the reaction occurs and had considerable strength after final set as seen in the compressive strength section. Nonetheless, the mortar specimens did not have enough strength to overcome the stresses induced due to the rapid change in temperature. Furthermore, since the cement paste reached 100 °C rapidly, water evaporated producing steam voids and probably tensile forces. The lack of strength to overcome the induced stresses can be related to this water phase change. Figure 7 shows that when 4.5 oz/100 lb. of cement (0.18%) is used, the temperature reaches 100 °C as well, but the cracks were not observed in these samples. In this case, since citric acid is added, this solute increases the boiling point of water and yields more time for it to react with cement [11].

Figure 10. Fracture of specimen after water curing.  
Source: Adapted from [24]

Figure 11. Cracks of specimen after air curing.  
Source: Adapted from [25]
E. CONCLUSIONS AND RECOMMENDATIONS

The following conclusions can be made based on the work described in this paper:

a. The relationship between setting time and citric acid dosage is linear. This has been confirmed experimentally based on Vicat testing, penetrometer tests, and temperature-time relationships from this research.

b. The results obtained from data available in this research and the modified methodology of the maturity method yielded a linear trend. Therefore, the fact that the equivalent age and citric acid dosage is linear supports the fact the setting time is behaves linearly with respect to citric dosage since the maturity method correlates setting time, reaction rate and equivalent age.

c. The strength of the BCSA specimens after setting and at one day is improved by increasing the dosage of citric acid until it reaches a breakpoint between 0.7% and 1.4% of dosage. At some dosage between 0.7% and 1.4% the after setting and 3 hours strength reduces significantly.

d. At later stages, the strength is slightly improved. A more uniform formation of ettringite due to a slower reaction and a lower porosity due to improved fluidity can be determined to be the main factors influencing this improvement in compressive strength.

e. The range of water temperatures used in this research did not have a significant effect on compressive strength.

f. Increasing the dosage of citric acid reduces the peak temperature and the temperature-time slope, but increases the area under the curve of the temperature-time plot.

g. The appropriate use of retarders, citric acid in this case, allows us to understand the behavior of BCSA cement paste under different dosage. Concrete should have the same behavior since the aggregates are not part of the reaction. However, additional research with BCSA concrete is required to understand its behavior and determine the possible positive outcomes of its use.

More research is required to fully understand its behavior and promote its usage in larger projects. This research paper offers some basic relationships between the dosage of citric acid and the cement behavior to be used to guide mixture proportioning and other research. Future work in this area could improve on the methodology presented here by making the following changes:
a. Variability in parameters such as setting time and compressive strength can be attributed to ambient temperature and humidity. Research with more control of these is required.
b. Curing in cold water is not recommended for very low dosage of citric acid. Even curing in air will required some special methodology to avoid fractures on the specimens.
c. Research with a wider water temperature range should be done to account for seasonal conditions in the field.
d. The use of other techniques such as calorimetry and X-ray diffraction can help to monitor the hydration rate and obtain other chemical information more precisely.
e. Applying the maturity method described by the ASTM C1074-17 is another alternative to confirm the facts about setting time presented in this paper.

F. REFERENCES

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G. APPENDIX

**Figure 12.** Penetrometer test plot at 21.1 °C water temperature.

**Figure 13.** Log-Log Penetrometer test plot at 21.1 °C water temperature.

**Figure 14.** Penetrometer test plot at 23.3 °C water temperature.
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Figure 15. Log-Log Penetrometer test plot at 23.3°C water temperature.

Figure 16. Penetrometer test plot at 25°C water temperature.

Figure 17. Log-Log Penetrometer test plot at 25°C water temperature.
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Figure 18. Vicat test plot for 36 oz dosage at 21.1 °C water temperature.

Figure 19. Vicat test plot for 36 oz dosage at 23.3 °C water temperature.

Figure 20. Vicat test plot for 36 oz dosage at 25 °C water temperature.
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Figure 21. Vicat test plot for 18 oz dosage at 21.1 °C water temperature.

Figure 22. Vicat test plot for 18 oz dosage at 23.3 °C water temperature.

Figure 23. Vicat test plot for 18 oz dosage at 25 °C water temperature.
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Figure 24. Vicat test plot for 13.5 oz dosage at 21.1 °C water temperature.

Figure 25. Vicat test plot for 13.5 oz dosage at 23.3 °C water temperature.

Figure 26. Vicat test plot for 13.5 oz dosage at 25 °C water temperature.
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Figure 27. Vicat test plot for 9 oz dosage at 21.1°C water temperature.

Figure 28. Vicat test plot for 9 oz dosage at 23.3°C water temperature.

Figure 29. Vicat test plot for 9 oz dosage at 25°C water temperature.
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Figure 30. Vicat test plot for 4.5 oz dosage at 21.1 °C water temperature.

Figure 31. Vicat test plot for 4.5 oz dosage at 23.3 °C water temperature.

Figure 32. Vicat test plot for 4.5 oz dosage at 25 °C water temperature.
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Figure 33. Vicat test plot for 0 oz dosage at 21.1 °C water temperature.

Figure 34. Vicat test plot for 0 oz dosage at 23.3 °C water temperature.

Figure 35. Vicat test plot for 0 oz dosage at 25 °C water temperature.
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```matlab
function TMS = Slopetemp(dosage)
    DS0oz = xlsread('Tempdata.xlsx',1,'AD5:AD33644');
    DS45oz = xlsread('Tempdata.xlsx',1,'AC5:AC50057');
    DS90oz = xlsread('Tempdata.xlsx',1,'AB5:AB69362');
    DS135oz = xlsread('Tempdata.xlsx',1,'AA5:AA92855');
    DS180oz = xlsread('Tempdata.xlsx',1,'Z5:Z100279');
    DS360oz = xlsread('Tempdata.xlsx',1,'AE5:AE210040');
    t_step = 0.1/60;
    if dosage == 0
        S = gradient(DS0oz(601:4501),t_step);
        TMS = t_step*(find(S==max(S))+601);
    elseif dosage == 4.5
        S = gradient(DS45oz(9001:15001),t_step);
        TMS = t_step*(find(S==max(S))+9001);
    elseif dosage == 9
        S = gradient(DS90oz(18001:30001),t_step);
        TMS = t_step*(find(S==max(S))+18001);
    elseif dosage == 13.5
        S = gradient(DS135oz(12001:39001),t_step);
        TMS = t_step*(find(S==max(S))+12001);
    elseif dosage == 18
        S = gradient(DS180oz(24001:60001),t_step);
        TMS = t_step*(find(S==max(S))+24001);
    elseif dosage == 36
        S = gradient(DS360oz(102001:123001),t_step);
        TMS = t_step*(find(S==max(S))+102001);
    end
```

Figure 36. MATLAB code used to determine location of maximum slope.
function AUCBP = AreaTemp(dosage)
    Timeshift = xlsread('Tempdata.xlsx',1,'Y5:Y210040');
    DS0oz = xlsread('Tempdata.xlsx',1,'AD5:AD33644');
    DS45oz = xlsread('Tempdata.xlsx',1,'AC5:AC50057');
    DS90oz = xlsread('Tempdata.xlsx',1,'AB5:AB69362');
    DS135oz = xlsread('Tempdata.xlsx',1,'AA5:AA92855');
    DS180oz = xlsread('Tempdata.xlsx',1,'Z5:Z100279');
    DS360oz = xlsread('Tempdata.xlsx',1,'AES:AE210040');
    if dosage == 0
        AUCBP = trapz(Timeshift(1:18455), DS0oz(1:18455));
    elseif dosage == 4.5
        AUCBP = trapz(Timeshift(1:28535), DS45oz(1:28535));
    elseif dosage == 9
        AUCBP = trapz(Timeshift(1:44297), DS90oz(1:44297));
    elseif dosage == 13.5
        AUCBP = trapz(Timeshift(1:62147), DS135oz(1:62147));
    elseif dosage == 18
        AUCBP = trapz(Timeshift(1:78101), DS180oz(1:78101));
    elseif dosage == 36
        AUCBP = trapz(Timeshift(1:161346), DS360oz(1:161346));
    end

Figure 37. MATLAB code used to determine the area under the curve of the temperature-time plots.