A comparison study of Portland cement hydration kinetics as measured by chemical shrinkage and isothermal calorimetry

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1. Introduction

The hydration of Portland cement is a complex process. Despite decades of research, many detailed features are still not clearly understood today, primarily because of the complicating influences of different clinker phases, impurities, and their interactions. Nevertheless, the general hydration kinetics of cement is often represented by the rate of change of the overall degree of hydration $\alpha$, which is defined as the total weight fraction of cement reacted. As a composite material consisting mainly of four compounds, or clinker phases (C3S, C2S, C3A and C4AF), the overall degree of hydration of cement is typically written as [1]:

$$\alpha(t) = p_{C_3S}x_{C_3S}(t) + p_{C_2S}x_{C_2S}(t) + p_{C_3A}x_{C_3A}(t) + p_{C_4AF}x_{C_4AF}(t)$$  (1)

where $p_i$ is the original weight fraction of Phase $i$ in the anhydrous cement and $x_i(t)$ is the degree of hydration of Phase $i$ at time $t$. Direct determination of $x_i(t)$ can be made by quantitative X-ray diffraction analysis (QXDA) [1,2], though it is difficult to obtain accurate results. Some properties of a hydrating cement paste, such as the non-evaporable water content, the cumulative heat evolution and the total chemical shrinkage have been shown to have approximately linear relationships with the overall degree of hydration [1.3–5]. As a matter of fact, $\alpha$ is more easily and commonly determined indirectly by tracking the time dependence of one or more of these properties. For the purposes of this paper, the semi-continuous measure of the progress of the degree of hydration with time ($\alpha(t)$) and its derivative ($d\alpha(t)/dt$) are both referred to as the hydration kinetics curves.

Among the different methods of approximating cement hydration progress, heat of hydration tests used to be the only one that provided continuous data suitable for evaluating hydration mechanisms. Automated chemical shrinkage test methods have been developed in recent years [6–9] and are now also frequently used to study cement hydration mechanisms [9–12]. Therefore, it is important to investigate the differences and similarities between these two different methods, which may also help us gain further insights about the cement hydration process. In this study, both the traditional isothermal calorimetry and one of the newly developed chemical shrinkage test methods were employed to measure the hydration kinetics of cement at similar curing conditions and a detailed comparison of the two sets of test results was performed.

The experimental test methods and detailed test plan of this study are described Section 2. The theoretical analysis regarding the correlation between heat of hydration and chemical shrinkage is presented in Section 3. Because temperature control of the chemical shrinkage tests performed in this study was not very accurate, a method of correcting the temperature differences between isothermal calorimetry tests and chemical shrinkage tests...
is also proposed in Section 3 based on a scale factor model developed earlier. The scale factor model was developed to simulate the effect of curing temperature and pressure on cement hydration kinetics and has been preliminarily validated using chemical shrinkage test data [12]. When the hydration kinetics curve of a reference curing temperature is known for a given cement paste, the scale factor model can be used to estimate the hydration kinetics curve for a given temperature or the temperature of a given hydration kinetics curve. In Section 4.1, the model is further validated with isothermal calorimetry test data from this study. Since the rate of hydration is measured directly in these tests, the accuracy and limitations of the scale factor model can be observed and discussed in greater detail. In Section 4.2, it is shown that the hydration kinetics measured by chemical shrinkage can be correlated excellently with those measured by isothermal calorimetry by applying the proposed model in this study.

2. Materials and methods

2.1. Materials

Oilwell cements are based upon Portland cement, but manufactured to a higher level of consistency from one production batch to another. There are no substantial differences between oilwell cements and ordinary Portland cements, except for the fact that some Class H (API Specification 10A [13]) cement may have near zero CaO content. API Classes A, B, and C cements are similar to ASTM Types I, II, and III cements, respectively. The hydration kinetics of four different classes of oilwell cements, namely Classes A, C, G, and H, were investigated at different curing temperatures in this study. All slurries (cement pastes) were prepared with de-aerated water and cement only, with no additives. As will be shown in Section 2.2, standard water-to-cement (w/c) mass ratios for each class of cement were used, as defined in API Specification 10A [13]. The main compound compositions of the different types of cements derived from the oxide analysis test results using the Bogue calculation method [14] are presented in Table 1. As shown in the table, two different types of Class H cements were used: a premium Class H (H-P) and a standard Class H. Some minor composition variations were observed between different batches of standard Class H cement (H-I and H-II) produced from the same plant.

The particle size distributions of the cements were measured by the laser scattering technique with dry dispersion methods (at least 10 measurements were performed on each type of cement). The average test results are presented in Fig. 1. Class H-I cement was not measured as it should be similar to Class H-II cement. The median particle sizes for Classes A, C, G, H-P, and H-II cements were 38 μm, 15 μm, 34 μm, 30 μm, and 23 μm, respectively, while their specific surface areas calculated from the PSD data (assuming spherical particles and a cement density of 3150 kg/m$^3$) were 356 m$^2$/kg, 565 m$^2$/kg, 327 m$^2$/kg, 394 m$^2$/kg, and 323 m$^2$/kg, respectively. Apparently, the median particle size does not necessarily correlate with the specific surface area since the latter is primarily dominated by the relative proportion of the fine particles. The particle size distribution curves for the Class A, G, and C cements are very similar, suggesting that similar grinding procedures may have been adopted in manufacturing these cements. Class C cement is ground much finer than the other classes to achieve a higher specific surface area and enhance early-age reactivity.

2.2. Chemical shrinkage test

The total absolute volume of cement hydration product is smaller than the combined initial volume of the anhydrous cement and water. This reduction in volume during hydration is known as chemical shrinkage. Despite the apparently simple principle of chemical shrinkage measurement, there are experimental difficulties that can yield spurious results. For example, the traditional chemical shrinkage test (ASTM C1608 [15]) involves adding a significant amount of water on top of a thin specimen (<10 mm) to keep it saturated. A recent study showed that the quantity and composition of the surface water have a significant impact on test results [16]. Increasing the amount of surface water was found to increase the chemical shrinkage rate before the end of the induction period and to reduce the peak chemical shrinkage rate during later periods. The initial increase is probably due to the accelerated dissolution as a result of more Ca$^{2+}$ leaching out of the paste sample (and into the surface water solution), while the reduced peak rate might be related to a dilution effect. Massive precipitates of portlandite were observed on the surface of specimens of alite (the main composition of Portland cement) used for chemical shrinkage measurement [17]. Additionally, when the same measuring device was used, increasing sample thickness was consistently found to cause a reduction in chemical shrinkage at later ages (>15 h) [6,16,17]. This thickness effect may be explained by two hypotheses: (1) the reduction in the permeability of the sample might prevent surface water from filling all the pores in the thicker samples (depercolation); (2) a larger fraction of the thinner sample is diluted by the surface water, resulting in a faster hydration rate at later ages. Costoya [17] found that using a small diameter device with less surface water (cylindrical flask) systematically gave a higher chemical shrinkage of a given mass of alite paste than when the same mass of paste was used in a large diameter device with more surface water (Erlenmeyer flask), even though the former generated a much thicker sample. The author also found that chemical shrinkage measured with the former device was the same as that measured with a set ground paste sample for a period exceeding 250 h. Therefore, for cement pastes with relatively high w/c ratios, surface water probably has a much stronger effect on the test result than the thickness of the sample and the latter is probably not a limiting factor.

![Fig. 1. Particle size distributions of different classes of cements.](image-url)
Table 2
Chemical shrinkage tests (test series I, test duration = 72 h).

<table>
<thead>
<tr>
<th>Cement temperature (°C)</th>
<th>Ambient (^a)</th>
<th>40° (^b)</th>
<th>60° (^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.46</td>
<td>CS-A-1</td>
<td>CS-A-2</td>
</tr>
<tr>
<td>C</td>
<td>0.56</td>
<td>CS-C-1</td>
<td>CS-C-2</td>
</tr>
<tr>
<td>G</td>
<td>0.44</td>
<td>CS-G-1</td>
<td>CS-G-2</td>
</tr>
<tr>
<td>H-P</td>
<td>0.38</td>
<td>CS-HP-1</td>
<td>CS-HP-2</td>
</tr>
<tr>
<td>H-I</td>
<td>0.38</td>
<td>CS-H-1</td>
<td>CS-H-2</td>
</tr>
</tbody>
</table>

\(^a\) Lab temperature (\(\sim 24 \pm 2.8\) °C).
\(^b\) Estimated specimen temperature (will be corrected later based on test results).

In this study, chemical shrinkage was measured by a recently developed test apparatus [18] originally designed to obtain in situ tensile strength of oilwell cements at different curing pressures. The apparatus allows automatic collection of chemical shrinkage data from samples cured at different temperatures and pressures. In addition, by using a large volume of cement paste (four 51 mm by 170 mm cylinders), the effect of surface water on test results is expected to be minimal. When a hydrostatic curing pressure of 0.69 MPa was applied, tests performed using hollow cylinders (whose entire annular surfaces were covered by filter paper for water saturation) with a wall thickness of approximately 10 mm were found to generate the same results as solid cylinders [18], suggesting sample thickness is not a limiting factor of test results for this type of test under the conditions being employed. The test design of the chemical shrinkage test series is shown in Table 2.

The test series was conducted at 40, 60, and 80 °C curing temperatures. The focus of this study is the correlation between cumulative heat evolution and hydration of clinker phases, respectively (typically in J/g cement); and CS(t) and CS\(^e\) are the total chemical shrinkage at time \(t\) and at complete hydration, respectively (typically in mL/g cement). Since \(\omega_0\) depends on the molar masses of the hydration products, while \(H^f\) depends on the enthalpy changes of the chemical reactions, both of which are expected to remain constant as long as the chemical formulae of the hydration products do not change. However, CS\(^e\) depends on the molar volumes of water and the hydration products and hence varies with both temperature and pressure. According to Eq. (2), since \(\omega_0\) and \(H^f\) remain invariant (at least within the temperature and pressure range used in this study), the dependency of CS\(^e\) on temperature and pressure can be approximately evaluated by studying the correlations between \(w_0\), \(H(t)\), and \(CS(t)\) at various curing conditions. Studies have shown that the ratio of CS\(^e\) to \(w_0\), evaluated at discrete data points, decreases with increasing temperature [6,10], suggesting that CS\(^e\) decreases with increasing temperature. The focus of this study is the correlation between CS\(^e\) and \(H(t)\), both of which have been measured continuously.

It should be pointed out that the indirect methods only give a gross approximation to the hydration rate of cement because hydration of the different phases also progresses at different rates. If we ignore the interactions between different clinker phases and the phase changes of different hydration products during hydration, then the total rate of hydration and chemical shrinkage may be correlated with the degree of hydration of each individual clinker phase as follows:

\[
\begin{align*}
H(t) &= a_{C5P1C5}x_{C5}(t) + a_{C3P1C3}x_{C3}(t) + a_{C3P1C3}x_{C3}(t) + a_{C3P1C3}x_{C3}(t) \\
&+ a_{C3P1C3}x_{C3}(t) + a_{C3P1C3}x_{C3}(t) + a_{C3P1C3}x_{C3}(t) + a_{C3P1C3}x_{C3}(t)
\end{align*}
\]

(3)

\[
\begin{align*}
CS(t) &= b_{C5P1C5}x_{C5}(t) + b_{C3P1C3}x_{C3}(t) + b_{C3P1C3}x_{C3}(t) + b_{C3P1C3}x_{C3}(t) \\
&+ b_{C3P1C3}x_{C3}(t) + b_{C3P1C3}x_{C3}(t) + b_{C3P1C3}x_{C3}(t) + b_{C3P1C3}x_{C3}(t)
\end{align*}
\]

(4)

where \(a_i\) and \(b_i\) are the cumulative heat evolution and the total chemical shrinkage, respectively, associated with the complete hydration of 1 g of clinker phase \(i\). Eqs. (3) and (4) can be used to estimate the parameters associated with the complete hydration condition in Eq. (2), using the sums

\[
H^f = \sum a_i p_i, CS^e = \sum b_i p_i
\]

(5)

3. Theoretical background and analysis

3.1. Indirect methods of measuring cement hydration

When the indirect methods are employed to measure cement hydration progress, the relationships between experimental results and the overall degree of hydration can be expressed as [9,21–25]:

\[
\alpha(t) = \frac{w_0(t)}{w_0^0} = \frac{H(t)}{H^f} = \frac{CS(t)}{CS^e}
\]

(2)

where \(w_0(t)\) and \(w_0^0\) are the non-evaporable water content at time \(t\) and at complete hydration, respectively (typically in g/g cement); \(H(t)\) and \(H^f\) are the cumulative heat evolution at time \(t\) and at complete hydration, respectively (typically in mL/g cement); and CS(t) and CS\(^e\) are the total chemical shrinkage at time \(t\) and at complete hydration, respectively (typically in mL/g cement). Since \(\omega_0\) depends on the molar masses of the hydration products, while \(H^f\) depends on the enthalpy changes of the chemical reactions, both of which are expected to remain constant as long as the chemical formulae of the hydration products do not change. However, CS\(^e\) depends on the molar volumes of water and the hydration products and hence varies with both temperature and pressure. According to Eq. (2), since \(\omega_0\) and \(H^f\) remain invariant (at least within the temperature and pressure range used in this study), the dependency of CS\(^e\) on temperature and pressure can be approximately evaluated by studying the correlations between \(w_0\), \(H(t)\), and \(CS(t)\) at various curing conditions. Studies have shown that the ratio of CS\(^e\) to \(w_0\), evaluated at discrete data points, decreases with increasing temperature [6,10], suggesting that CS\(^e\) decreases with increasing temperature. The focus of this study is the correlation between CS\(^e\) and \(H(t)\), both of which have been measured continuously.

2.3. Isothermal calorimetry test

Compared to chemical shrinkage, isothermal calorimetry is a more established test method of measuring overall cement hydration progress. In the second test series, hydration of the different types of cements is tracked with an isothermal calorimeter according to standard test procedures [19]. Tests were conducted at atmospheric pressure and three different curing temperatures. Table 3 shows the test design for this test series. The temperatures of isothermal calorimetry tests can be controlled more precisely due to the small sample size (4–5 g). For this technique, the average absolute difference between replicate specimens of cement paste is 2.4 × 10^{-5} W/g (cement), with a maximum absolute difference of 0.00011 W/g (cement), for measurements conducted between 1 h and 7 d after mixing [20]. It should be mentioned that the samples used in the isothermal calorimetry tests were cured under sealed condition, which is different from the saturated curing condition for the chemical shrinkage tests. Although it is difficult to accurately evaluate the effect of saturation condition on the isothermal calorimetry test results due to various experimental difficulties, a previous study [5] suggests that saturation has a negligible effect on tests with relatively high w/c ratios (the small differences in tests results were probably caused by the surface water effect discussed in Section 2.2).

Table 3
Isothermal calorimetry tests (test series II, test duration = 168 h).

<table>
<thead>
<tr>
<th>Cement temperature (°C)</th>
<th>25</th>
<th>40</th>
<th>60</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.46</td>
<td>IC-A-1</td>
<td>IC-A-2</td>
</tr>
<tr>
<td>C</td>
<td>0.56</td>
<td>IC-C-1</td>
<td>IC-C-2</td>
</tr>
<tr>
<td>G</td>
<td>0.44</td>
<td>IC-G-1</td>
<td>IC-G-2</td>
</tr>
<tr>
<td>H-P</td>
<td>0.38</td>
<td>IC-HP-1</td>
<td>IC-HP-2</td>
</tr>
<tr>
<td>H-I</td>
<td>0.38</td>
<td>IC-H-1</td>
<td>IC-H-2</td>
</tr>
</tbody>
</table>
The scale factor associated with temperature change from $T_i$ to $T$ and pressure change from $P_i$ to $P$.

One of the most important advantages of the scale factor model is that it is very straightforward and easy to use. The physical meaning of the model is that the hydration rate at any curing condition is increased or decreased by a factor of $C$ compared with that at the reference curing condition at the same degree of hydration.

Probably due to the different mechanism of hydration during the very early period (before the end of the induction period), it is sometimes necessary to offset the predicted hydration kinetics curve using the scale factor model to generate a better agreement with the experimental curve, especially when their curing temperatures are different. The offset is not necessary for tests performed at different curing pressures in the range from 0.7 MPa to 51.7 MPa [12]. Therefore, a more accurate representation of the relationship between the hydration kinetics curves at temperatures $T$ and $T_0$ (for the same curing pressure) is,

$$C_{T_0-T} = \exp \left( \frac{E_a}{R \left( T_0 - T \right)} \right)$$

where $a_T$ and $a_{T_0}$ are the functions representing the time dependence of the degree of hydration at temperatures $T$ and $T_0$, respectively; $t_0$ is the offset time, which appears to increase with increasing differences between $T$ and $T_0$.

4. Test results and discussion

4.1. Validation of the scale factor model

One of the most important assumptions of the scale factor model is that the normalized rate of hydration versus degree of hydration

<table>
<thead>
<tr>
<th>Phase</th>
<th>$C_5S$</th>
<th>$C_3S$</th>
<th>$C_3A$</th>
<th>$C_4AF$</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a_0$ (J/g)$^2$</td>
<td>510</td>
<td>247</td>
<td>1356</td>
<td>427</td>
<td>[28]</td>
</tr>
<tr>
<td>$b_0$ (ml/g at 25°C)</td>
<td>0.0596</td>
<td>0.0505</td>
<td>0.13</td>
<td>0.0409</td>
<td>[9]</td>
</tr>
<tr>
<td>$a_1$</td>
<td>8557</td>
<td>4011</td>
<td>10431</td>
<td>9104</td>
<td></td>
</tr>
</tbody>
</table>

* Obtained by multi-linear regression analysis from experimental data of 21 different cements, slightly different from theoretical values calculated from standard enthalpy of formation [28,29], which depend on the chemical formulae of hydration products.

If we consider an isothermal calorimetry test performed at temperature $T_0$ and a chemical shrinkage test performed at temperature $T_{CS}$, the degree of hydration measured by the former can be expressed as,

$$x_{T_0}(t) = \frac{H_{T_0}(t)}{H^0}$$

while that measured by the latter can be expressed as,

$$x_{T_{CS}}(t) = \frac{CS_{T_{CS}}(t)}{CS(T_{CS})}$$

Note that, as already discussed, $CS(T)$ is a function of temperature. Employing the scale factor model, one can obtain,

$$x_{T_{CS}}(t) = x_{T_0} C_{T_0-T_{CS}} (t - t_0)$$

where the scale factor is related to the temperature difference between the two types of tests by the following equation,

$$C_{T_0-T_{CS}} = \exp \left( \frac{E_a}{R \left( T_{CS} - T_0 \right)} \right)$$

By combining Eqs. (13)-(15), the cumulative heat evolution $H(t)$ can be related to the total chemical shrinkage $CS(t)$ by the following equation,

$$H_{T_{CS}}(t) = \frac{H^0}{CS(T_{CS})} CS_{T_{CS}}(C_{T_0-T_{CS}}(t - t_0))$$

### Table 4

Coefficients for estimating the parameters at the complete hydration condition.

<table>
<thead>
<tr>
<th>Phase</th>
<th>$C_5S$</th>
<th>$C_3S$</th>
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</tr>
<tr>
<td>$a_1$</td>
<td>8557</td>
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<td>10431</td>
<td>9104</td>
<td></td>
</tr>
</tbody>
</table>

* Obtained by multi-linear regression analysis from experimental data of 21 different cements, slightly different from theoretical values calculated from standard enthalpy of formation [28,29], which depend on the chemical formulae of hydration products.
The curve of a given cement paste is invariant with curing condition (for isothermal and isobaric tests). The assumption has been verified for different types of cement in the pressure range from 0.7 to 51.7 MPa using chemical shrinkage test data [12]. Oscillations of chemical shrinkage test results at high temperatures made it difficult to derive the rate of hydration accurately. Therefore, further validation of this assumption is desirable for different curing temperatures with the heat evolution data. Based on the values listed in Table 4 and assuming $C_2F$ generates the same amount of heat as $C_4AF$ on the same mass basis, the cumulative heat generated at complete hydration ($H_0$) was estimated to be 497.7 J/g (cement), 461.1 J/g, 470.1 J/g, 429.2 J/g, and 385.9 J/g, for Classes A, C, G, H-I, and H-P cements, respectively. The hydration progress of these different types of cement can be obtained by normalizing the heat evolution data by their respective $H_0$ values.

It should be pointed out that the scale factor model was developed for a single reaction process assuming that the curing condition only changes the rate of the reaction, but not its nature. In reality, Portland cement hydration is a much more complex process with all the different clinker phases having different reaction rates as well as different sensitivities to curing temperature changes (i.e., activation energies). The composition of the hydration products may also change if curing temperature changes significantly. Therefore, the model can be applied only approximately to analyze Portland cement hydration. Fig. 2 shows the rate of hydration vs. degree of hydration derived from the heat evolution data of Class H-I cement before and after normalization. While the normalized data at different curing temperatures coincide relatively well during early and late periods, slight deviations are observed during the middle period, (i.e., for degrees of hydration approximately ranging from 0.2 to 0.5). Fig. 3 shows the normalized rate of hydration vs. degree of hydration of the four other types of cement obtained at different curing temperatures. Similar convergence behaviors were observed except for the Class A and Class C cements at 60 °C, both of which diverge significantly from their behavior at 25 °C and 40 °C. For all the different types of cement, the divergences of the normalized kinetics curves becomes more significant as the temperature difference increases, due to the different temperature sensitivities of different phases. The normalized hydration data at 25 °C and 40 °C appear to show better
compliance with the model, indicating that the scale factor model is more accurate for relatively small temperature changes.

According to Eq. (11), the scale factor relating the hydration rates at two different curing temperatures can be estimated by applying basic coordinate transformation rules to transform the hydration kinetics curve at $T_r$ to achieve the best agreement with that at $T$ (the method is referred to as the best fit method). Fig. 4 shows the results of transforming the experimental hydration data obtained at 25 °C and 40 °C to fit/predict those at higher curing temperatures for Class C cement. Although it seems impossible to achieve perfect agreement for the entire curing period, the predictions are in good agreements with the actual experimental results during the period up to the first peak, which is mainly associated with $C_3S$ hydration. The second peak probably includes the contribution from $C_3A$ hydration, which reportedly has a higher activation energy than the overall value for Portland cement [31,32]. Therefore, the transformed hydration data from lower temperatures typically underestimate the second peak (Fig. 4). Applying a different scale factor for each individual phase in Portland cement would perhaps generate a better fit. Unfortunately, it is not yet possible to accurately and reliably measure the hydration progress of the different phases separately.

Fig. 5 shows the measured and predicted hydration kinetics curves for the other types of cement used in this study. The predictions are most accurate for the Class H-I cement, which has almost no $C_3A$, and least accurate for the Class A cement, which has the highest $C_3A$ content. This further confirms the hypothesis that a higher $C_3A$ activation energy can skew the model results. It is also interesting to note that the transformed data from lower temperatures typically overestimate slightly the degree of hydration at
later ages, with the exception of Class H-P cement, for which the opposite is true. Table 5 lists the offset time and scale factor, as well as the apparent activation energy (calculated from the scale factor using Eq. (12)) associated with each temperature change. The constants are obtained by trial and error to provide the best agreements as shown in Figs. 4 and 5. For comparison purposes, Table 5 also lists the scale factors and apparent activation energies calculated with the peak hydration rate method as proposed in a previous study [12]. The constants obtained by the two different methods generally agree well with each other for relatively small temperature changes (from 25 °C to 40 °C and from 40 °C to 60 °C) but may differ quite noticeably for the larger temperature change (from 25 °C to 60 °C). The apparent activation energies obtained in different temperature ranges appear to decrease with increasing temperature except for Class A cement, for which the opposite is true. The apparent activation energies obtained in this study are also found to be much lower than those calculated from chemical shrinkage test data [12]. The latter are probably less accurate due to the inadequate temperature control of the tests and errors associated with estimating CS³ at different temperatures (T⁹ is assumed to be independent of temperature).

4.2. Correlations between chemical shrinkage and heat of hydration of cement

Just as the hydration kinetics curve at a reference temperature can be transformed to fit the curves obtained at other temperatures, the chemical shrinkage curve can be transformed to fit the heat evolution curve (and vice versa) according to their correlations developed in Section 3.3. Following Eq. (17), the correlation factor $H^{0}$/CS³(TCS), the scale factor $C_{r-T}$, and the offset time $t_{0}$, can all be estimated by transforming the chemical shrinkage curve to achieve the best agreement with the heat evolution curve. Since the scale factor model is more accurate for smaller temperature changes, the temperature of the chemical shrinkage test should be close to that of the isothermal calorimetry test to obtain more reliable estimates. Fig. 6 shows the heat evolution curves of the Class H-P cement measured at 25 °C and the transformed chemical shrinkage curves (measured at lab temperature) that exhibit the hydration rate measured by chemical shrinkage seems to be higher than that measured by heat evolution.

Transpared heat evolution curves vs. transformed chemical shrinkage curves (Class H-P cement, w/c = 0.38, 25 °C).

The integral curves of heat evolution of different cements cured for 40–60 h are shown in Fig. 7. The heat evolution rate curves of the other types of cement measured at 25 °C and their corresponding best-fit transformed chemical shrinkage curves. Similar to the Class H-P cement, excellent agreement is obtained between the heat evolution curve and the transformed chemical shrinkage curve for the Class H-I cement. However, for cements that contain significant C₃A, the agreements between the two types of curves are somewhat poorer, especially around the main hydration peaks. The hydration of C₃A first produces ettringite during the hydration peak, which then further reacts with excess C₃A and transforms to calcium monosulfaloaluminate at later ages [26,28]. The test results show that the second peak rate (probably caused by C₃A hydration) measured by chemical shrinkage is always higher than that measured by heat evolution and that the rate measured by the former also decreases faster during the deceleration stage. Therefore, the ratio of chemical shrinkage to heat evolution associated with C₃A hydration is higher than that associated with C₃S hydration during the initial reaction (to form ettringite) and probably the reverse is true during the later reaction (to form monosulfaloaluminate). It should be noted that the ratio of $a₂$ to $b₂$ associated with C₃S hydration as shown in Table 4 represents the long-term results of some average cements and is not strictly applicable here.

The integral curves of heat evolution of different cements cured at different temperatures are compared with the transformed chemical shrinkage curves (measured at similar temperatures) in Fig. 8. In general, excellent agreements can be obtained between

<table>
<thead>
<tr>
<th>Table 5</th>
</tr>
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<tbody>
<tr>
<td><strong>Activation energies obtained from different analysis methods.</strong></td>
</tr>
<tr>
<td>Cement</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>A</td>
</tr>
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<td></td>
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<tr>
<td></td>
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<tr>
<td>C</td>
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<tr>
<td>G</td>
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<td>H-P</td>
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<tr>
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the integral hydration kinetics curves obtained from the two different methods. The constants used to transform the chemical shrinkage curves to fit the heat evolution curves, including the correlation factor, the scale factor, and the offset time, are listed in Tables 1 and 2.

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**Fig. 7.** Heat evolution vs. transformed chemical shrinkage for the different types of cement (derivative curves).

**Fig. 8.** Heat evolution curves vs. transformed chemical shrinkage curves for different types of cement at different curing temperatures (integral curves).
The cause of the offset time (primarily observed for ambient temperature tests) is not yet clear. It may be associated with variations of the duration of the induction periods, which tend to be affected by many different factors that are difficult to control. Following Eq. (16), the temperature of the chemical shrinkage test ($T_{CS}$) can be estimated from the more-precisely-measured temperature of the isothermal calorimetry test ($T_{IC}$) by using the scale factor and the previously obtained activation energies in the 25 °C to 60 °C temperature range (Table 5). As shown in Table 6, the calculated sample temperatures of the chemical shrinkage tests are slightly higher than the previously estimated values (Table 3). To further demonstrate that the scale factors are indeed associated with the slight temperature differences between chemical shrinkage tests and isothermal calorimetry tests, one of the chemical shrinkage tests (test CS-H-2) was performed by reducing the target temperature (set with the temperature controller) by 2.8 °C (5 °F). The obtained chemical shrinkage curve of this particular test is found to be directly proportional to the heat evolution curve (i.e., $C_0 = 1$).

At lab temperature (approximately 25 °C), the correlation factor between heat evolution and chemical shrinkage ($\frac{H^0}{C^0}$) for the different cements is found to range from 7500 J/mL to 8000 J/mL, well within the previously reported range of 6500–8500 J/mL [33,34]. The correlation factor increases with increasing curing temperature. Since $H^0$ is independent of curing temperature, the results suggest that $C^0$ increases with increasing temperature, consistent with previous studies [6,10]. In the relatively small temperature range investigated (25–60 °C), the variations of $C^0$ and $H^0$ with temperature can both be approximated by linear models. Fig. 9 shows the dependence of $C^0$ on curing temperature for different types of cement. The least square fits of all data points indicate the average rate of change to be $-0.00076$ mL/kJ per °C for $C^0$ and $+58$ J/mL per °C for $H^0$. When normalized to the values at 25 °C, the linear reduction rate of $C^0$ with increasing temperature is determined to be 0.57%, 0.65%, 0.58%, 0.66%, and 0.57% per °C for Class A, C, G, H-P, and H-I cements, respectively, suggesting an average reduction rate of 0.60% (with a standard deviation of 0.046%) per °C. For comparison, a reduction rate of 0.78% per °C may be obtained from test data of Zhang et al. [10] in the same temperature range, where the correlations between $w_0$ and $C^0$ of Class H cement were investigated.

5. Conclusions

The hydration kinetics of different types of cement cured at different temperatures has been evaluated by both the traditional isothermal calorimetry test method and a newly developed chemical shrinkage test method. A scale factor model that can be used to predict the effect of curing temperature on cement hydration kinetics, previously validated with chemical shrinkage test data, is further discussed and validated with the heat evolution test data. Due to the fact that the different compounds in cement hydrate at different rates and have different temperature sensitivities, the model is found to be more accurate for cement with simpler compositions (e.g., no or very low C3A content) and/or for smaller temperature changes (e.g. ≤15 °C). The scale factor model also introduces a new method of estimating the apparent activation energies of Portland cement (i.e. the best-fit method). The apparent activation energies of Classes A, C, G, H-P and H-I cements determined using the heat of hydration data in the temperature range of 25–60 °C are 40.2 kJ/mol, 37 kJ/mol, 40.2 kJ/mol, 35.5 kJ/mol, and 37 kJ/mol, respectively.

A correlation study of chemical shrinkage and isothermal calorimetry test data indicate that total chemical shrinkage is proportional to cumulative heat evolution if the two tests are conducted on the same cement paste at the same curing temperature, in agreement with previous research. The scale factor model discussed in this study can be used to account for any unmeasured small temperature differences. The proportionality constant, namely the ratio of total chemical shrinkage to total heat release at complete hydration ($C^0$/$H^0$), varies slightly with cement composition and decreases with increasing curing temperature. If $H^0$ is assumed to be independent of curing temperature, then $C^0$ decreases approximately linearly with increasing temperature at a rate of 0.600 ± 0.046% per °C from the reference values at 25 °C for the different cements used in this study. Compared to C5 hydration, early C3A hydration (typically at the main hydration peak) seems to create a higher ratio of chemical shrinkage to heat evolution, while later C3A hydration (typically during the deceleration period) appears to create a lower ratio. As a result, the rate of chemical shrinkage curve typically has a slightly different shape than the heat flow curve for cements that contain significant C3A.

Acknowledgements

The continuous support of this study from Dr. Lewis Norman and Dr. Ron Morgan of Halliburton is cordially appreciated. We thank Mr. David Meadows of Halliburton for his help in developing the chemical shrinkage test apparatus and Ms. Donna Chen of Columbia University for her help in performing part of the experimental work. The critical review of this work by Dr. Jeffrey Bullard of NIST is also greatly acknowledged.

### Table 6

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<th>Test no.</th>
<th>$t_0$ (h)</th>
<th>$C_0$</th>
<th>$H^0/C^0$ ($T_{CS}$)</th>
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References


