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## **EVALUATING THE HYDRATION OF HIGH VOLUME FLY ASH MIXTURES USING CHEMICALLY INERT FILLERS**

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#### **Abstract**

Fly ash is frequently used as a replacement for cement in concrete. However, questions remain regarding the influence that fly ash has on the hydration of cement. This paper examines physical aspects (e.g., surface nucleation, cement particle spacing) and chemical aspects (e.g., pozzolanic and hydraulic reactions) of the fly ash and cement in mixtures containing high volumes of fly ash. In addition to using fly ash, a chemically inert filler was used consisting of a blend of fine silica sands with approximately the same particle size distribution as that of the fly ash. The paper compares reactivity results from 1) cement, 2) cement-fly ash and 3) cement-inert filler systems. Isothermal calorimetry measurements are used to quantitively evaluate the role played by the fly ash in hydration of high volume fly ash mixtures. The results provide a decoupling of the physical and chemical effects of high volume fly ash on cement hydration.

#### **Keywords**

Filler effect; high volume fly ash; hydration; inert filler; isothermal calorimetry

#### **1. INTRODUCTION**

Fly ash is a by-product of coal combustion that has been broadly used by the concrete industry as a supplementary cementitious material (SCM) [1–9]. It can be used to replace cement, which decreases the clinker factor and embodied  $CO<sub>2</sub>$  [10–12] and generally improves workability and durability [13]. High Volume Fly Ash (HVFA) mixtures are

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typically designed with more than 50 % (by mass) of the cement replaced with fly ash. However, fly ash is less reactive than cement, which can sometimes cause retardation and extended setting time issues, especially when employed at higher volumes [14–16]. The degree of reactivity for the fly ash depends on the type of fly ash used (e.g., a Class C fly ash has typically more hydraulic properties than a Class F fly ash) [17]. With both classes of fly ash, lower strengths are typically observed in HVFA concrete at early ages [18]. Typically, it is possible to counteract this strength loss by reducing the water-to-cementitious ratio [19], using accelerating additions, replacing a portion of the fly ash with a fine limestone powder, or switching to a more reactive Type III cement [20–24].

The reactivity of fly ash and its effect on cement hydration has been a focus of study throughout the  $20<sup>th</sup>$  and into the  $21<sup>st</sup>$  century [19–23]. It now becomes more relevant to understand due to the current tendency of using larger amounts of fly ash to replace cement in concrete. However, the reaction kinetics of fly ash-cement systems are complicated by the fact that the reactions of the cement and fly ash may interact and, more importantly, by the difficulty in measuring the degree of reaction of these two components independently [25]. Recently, several studies have focused on developing faster and more reliable techniques for physical and chemical characterization of fly ash that can be related to fly ash reactivity [26–32].

According to ASTM C595, a pozzolan is defined as "a siliceous or siliceous and aluminous material, which in itself possesses little or no cementitious value but will, in finely divided form and in the presence of moisture, chemically react with calcium hydroxide to form compounds possessing cementitious properties (pozzolanic activity)." Fly ash is one of the materials that falls into this category. The pozzolanic reaction typically starts at later ages, since fly ash needs  $Ca(OH)_2$  formed during cement hydration for the reaction. Depending on the fly ash characteristics and the alkalinity of the pore solution, its reaction with  $Ca(OH)_2$ will occur earlier or later [33,34]. It is often thought that the pozzolanic effect is more dominant than any other effects (e.g., filler effects) [35–37], especially at later ages.

It is generally accepted that, under most circumstances, concrete containing fly ash is more durable than conventional ordinary portland cement (OPC) concrete, due mainly to the pore refinement produced by the pozzolanic reaction that converts the calcium hydroxide  $(Ca(OH)_2)$  formed during cement hydration to additional calcium silicate hydrate gel (C-S-H), resulting in a reduction in the permeability/diffusivity of the matrix [34,38]. The pozzolanic reaction can also cause a size reduction and densification of the interfacial transition zone (ITZ) region between the aggregates and the cement matrix, often considered to be a weak interface due to the higher porosity as compared to the bulk matrix. The reaction between fly ash and the  $(Ca(OH)<sub>2</sub>)$  in this ITZ region reduces its local porosity.

A common technique to determine the reactivity of cementitious materials is by using an isothermal calorimeter. The hydration reaction is exothermic; therefore, measuring the heat release of these materials indicates how much of the material has reacted. When comparing the heat release between plain cement and fly ash-cement systems, it can be mistakenly interpreted that the difference in heat release between these two systems is solely due to the fly ash reactivity. Typical cumulative heat release curves for cement and cement-fly ash

pastes are shown in Figure 1 where the heat release per g of cement of a 100 % OPC and a 40 % (by volume) fly ash  $-60$  % OPC system are plotted as a function of time. In the conventional practice of normalizing the heat release per g of cement (as in Figure 1), the fly ash is considered to act as an inert material; therefore, the difference in heat observed is purely due to the presence of fly ash, which can have physical and/or chemical effects on the cement hydration, instead of just providing additional chemical reactions (hydraulic and pozzolanic) that increase the heat release. Inert fillers with similar particle size distributions (PSD) to the pozzolanic materials under study have been used in the past in an attempt to decouple the chemical and physical effects of SCMs, based on compressive strength measurements [39].

The present investigation explores the idea that the chemical reactivity of the fly ash is not the only aspect to consider when fly ash is mixed with cement. Fly ash can also act like a filler [19], especially at very early ages, prior to its chemical reaction. In addition, retardation of the initial hydration reactions of cement (and consequently, of initial setting time) due to dilution and interaction with some fly ash components is commonly observed [21,40]. These effects have been examined in detail in systems containing fillers other than fly ash [41–43].

As mentioned above, the reactivity of the fly ash, particularly Class F fly ash, depends on the alkalinity of the pore solution and the availability of  $Ca(OH)_2$ , both of which build up over a few days. Therefore, the amount of reaction of fly ash in the first day or so is often negligible and changes in hydration kinetics may be dominated by the physical filler effect. Three combined physical effects may occur; 1) cement dilution as an increase in the watercement mass ratio (w/c) produces a larger separation distance between cement particles, 2) deflocculation (that is, a decrease in the flocculation level of cement particles, as the fly ash particles break up some of the flocculated cement 3-D network structure, also improving the paste rheology [44]), thus promoting a better cement hydration since more cement surfaces per unit volume of cement are exposed, and 3) provision of new nucleation sites via the fly ash surfaces that may promote the formation of additional hydration products. A graphical representation of these concepts is shown in Figure 2. This was done by adapting a hard core-soft shell (HCSS) model developed at NIST [40,45]. In this case, the model was used to represent a system where cement particles (red color) of different sizes (according to the particle size distribution of the cement used in this study) are randomly distributed in a three-dimensional volume. The same particle size distribution was used where 60 % (by volume) of the cement particles were replaced with either fly ash or inert filler (blue color). The image on the left represents a water-to-cement ratio,  $w/c = 0.30$  plain system with just cement particles, whereas the image in the center corresponds to the same system, but with 60 % of the cement particles (by volume) being replaced with fly ash (or inert filler), thereby increasing the number of potential nucleation sites and reducing the number of contacts between cement particles (deflocculation). Finally, the image on the right shows a system where the fly ash (or inert filler) particles are removed, thus increasing the cement particles' spacing (which would correspond to a  $w/c = 0.67$  plain system).

Because the heat flow and heat release plots are conventionally normalized by the mass of cement, the dilution effect typically only manifests itself at later ages. Conversely, at

early ages (to 48 h), plots for heat flow and heat release normalized by mass of cement are basically identical for  $w/c$  in the range of 0.3 to 0.5, for pastes prepared under high shear conditions [46,47]. At later ages, the calorimetry curves will diverge as the lower  $w/c$  pastes undergo a more significant slowing of their hydration than the higher  $w/c$  ones, due to space (porosity) limitations and self-desiccation [46,48].

#### **2. OBJECTIVE**

The objective of this study is to better understand the chemical contribution of fly ash as a reactive element (i.e., its pozzolanic or chemical activity) and its physical contributions as a nucleation agent and by increasing the cement particle spacing and reducing flocculation. Therefore, three different aspects will be analyzed:

- **1.** Evaluate the physical and chemical effect that high levels of fly ash replacement have on the cement hydration. This would include the aspects of pozzolanic reaction, additional nucleation sites/deflocculation, and increased cement particle spacing.
- **2.** Isolate the physical effects (nucleation/deflocculation and cement particle spacing) from the chemical effect (pozzolanic reaction).
- **3.** Differentiate, within the physical effect, between the contribution of fly ash on the provision of nucleation sites/exposed cement surface area and increased cement particle spacing.

#### **3. MATERIALS**

An ASTM C150-16e1 Type I/II OPC was used in this study, with a Blaine fineness of 476 m<sup>2</sup>/kg, a density of 3170 kg/m<sup>3</sup>  $\pm$  10 kg/m<sup>3</sup>, an estimated Bogue potential phase composition of 52 % C<sub>3</sub>S, 18 % C<sub>2</sub>S, 8 % C<sub>3</sub>A, and 9 % C<sub>4</sub>AF by mass, and a Na<sub>2</sub>O equivalent of 0.5 % by mass. A Class C fly ash (ASTM C618-15) was also used with a density of 2630 kg/m<sup>3</sup>  $\pm$  10 kg/m<sup>3</sup>. Silica sand with an apparent specific gravity of 2.65 was used as the fine aggregate. The chemically inert (hereafter designated as inert) filler used to replace the fly ash consisted of a combination of two size fractions of fine silica sand (SIL-5 and SIL-40) with a very similar density to that of the fly ash used,  $2650 \text{ kg/m}^3 \pm 10 \text{ kg/m}^3$ . (Here it is assumed that even the smallest particles of this material are chemically inert). Table 1 shows the chemical compositions of the materials used in the study.

Figure 3 shows the particle size distribution (PSD) of the fly ash used in the study. Two gradations of fine silica sand were chosen to comprise the inert filler. The two gradations selected (SIL-5 and SIL-40) were combined to have a similar PSD to that of the fly ash as shown in Figure 3. The mass proportions used in the combination were 16.7 % and 83.3 % of the SIL-5 and SIL-40, respectively. The result shown for fly ash is the average of six individual measurements and the error bars (one standard deviation) would fall within the size of the shown symbols.

It is important to note at this point that although the PSD obtained when combining two fractions of inert filler is similar, it is not identical to that of the fly ash. Also, the angular

shape of the inert filler (due to grinding) will be different from the generally round shape of the fly ash particles, potentially providing a different amount of nucleation sites (surfaces). As a reference point, for ground cement particles, their surface area is increased by about a factor of 1.4 relative to spherical particles of an equivalent volume [49].

#### **4. MIXTURE PROPORTIONS**

A total of nine mortar mixtures with 55 % of fine aggregate volume fraction were prepared. All mortars were prepared in a Hobart blender according to the mortar mixing procedure described in ASTM C305 [50]. The mixture proportions are shown in Table 2. The term  $w/c$  is used when the cementing phase is just cement, whereas  $w/cm$  is used when either fly ash or inert silica sand are used in the cementing fraction.  $V_w$  (paste) indicates the volume fraction of water in the paste portion of each mortar. It is not exactly constant for mortars M1 to M7 due to the decision to maintain  $w/cm$  constant and the differing densities of the cement, fly ash, and filler. As no high range water-reducing admixtures were employed in any of the mixtures, it is expected that the cement particles will be flocculated to the highest degree possible in each mixture. The reasons for choosing each of the mixture proportions are presented below:

- **•** Mixtures prepared to accomplish objective #1. Four mortars were prepared with  $w/c = 0.3$  and w/cm = 0.3: a plain mortar (M1), and three mortars containing different fly ash volume replacements (M2, M3, M4), with the fly ash replacing the volume of cement in M1 at volume replacement levels of 40 %, 60 %, and 80 %, respectively.
- **•** Mixtures to accomplish objective #2: Three mortars (M5, M6, and M7) were prepared with  $w/cm = 0.3$ . In these mortars, an inert filler (IF) is used to replace the cement in the same proportions as in M2, M3, and M4. The use of the inert filler corresponds to an equivalent system to that obtained in the fly ash mortars, without providing a chemical effect from the filler, thus isolating its physical action (i.e., M2 is similar to M5).
- Mixtures to accomplish objective #3: Two mortars were prepared with  $w/c$  of 0.47 and 0.67 (with neither inert filler nor fly ash). These  $w/c$  correspond to the same w/c used in mortars M2 and M3 (and therefore, M5 and M6), respectively. These systems would correspond to a similar cement particle spacing distribution in the matrix, while providing neither additional nucleation sites nor deflocculation, in other words, just considering the cement particle spacing. These mortars are labeled as M8 and M9. (NOTE: An attempt to reproduce the cement particle spacing of the mortar containing 80 % of either fly ash or inert filler was performed. This mortar mixture would have had a  $w/c$ = 1.30, but showed severe segregation/bleeding issues. Therefore, it was decided by the authors to not include this mixture in the analysis).

#### **5. EXPERIMENTAL**

Measurement of the heat release due to the exothermic reaction of cementitious materials can be used to assess the reaction of a cementitious system. Differences in heat release can be used to quantify the influence of fly ash or inert fillers. Approximately 5 g of the externally mixed mortar mixtures shown in Table 2 were weighed and placed in a glass ampoule, which was then capped/sealed and placed into an isothermal calorimeter about 10 min after cement was first mixed with water and then monitored for up to 7 d. The heat release and rate of heat release were determined. Previously, the average absolute difference between replicate specimens was measured to be  $2.4 \times 10^{-5}$  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 [44].

#### **6. RESULTS AND DISCUSSION**

Figures 5 and 6 show the measured heat release and heat flow per gram of cement for the control mixture ( $w/c = 0.3$  plain) and the mixtures containing 40 % by volume of either fly ash or inert filler. The plain mixture with  $w/c = 0.47$  is also included since it represents the same cement particle spacing as those having a volume replacement level of 40 %. As mentioned before, normalizing the heat release per gram of cement would result in all the plots being similar if the fly ash is inert. One aspect to point out is that the low  $w/c$ mixture (M1) could undergo significant self-desiccation at later ages, which might bias the comparison with the other three mixtures that are expected to exhibit a lower amount of self-desiccation. As such, a layer of water was carefully added on top of the M1 specimen to minimize self-desiccation, and without altering its  $w/c$ . Figure 5 shows differences in the heat release among the four mixtures. The magnitudes of these differences are indicated with arrows in the figure and preliminarily attributed to different effects (physical and/or chemical). For reference, the single operator uncertainty for 7 d cumulative heat release provided in ASTM C1702 [51] is 3.6 % and would correspond to about 10 J/g cement at 7 d for the data in Figure 5.

In Figure 7, the total amount of heat release by the fly ash mixture has been separated into the three effects indicated in Figure 5: increased spacing, nucleation (including deflocculation) and fly ash reaction. These would have to be added to the heat released by the  $w/c = 0.3$  mixture (M1) to add up to the total represented by the fly ash mixture (M2). This was done by subtracting the amount of heat release among the curves. For instance, the heat attributed to spacing was calculated by subtracting the heat of M1 from the heat of M8. In Figure 7, the curves of each effect are presented in a stacked area array to visualize the gap filling between the M1 and M2 curves.

When comparing specimens M1 and M8 from Figure 5, a higher heat release is achieved in M8, being 8 % larger at 7 d. This is attributed to the increased cement particle spacing that promotes a further hydration. While there is practically no difference in the cumulative heat release during the first 24 h between M1 and M8 (in agreement with previous results [50, 53] and as should be expected since the samples maintain high humidity at very early ages), the results in Figure 6 do show that the lower  $w/c$  mixture (M1) presents greater heat

flow compared to M8 between 4 h and 10 h. The same effect was also observed by Castro et al. [48]. This might be attributed to the mixing intensity employed in this study. Bentz et al. [46] observed nearly identical heat release curves for cement pastes of different w/c prepared in a high shear blender; however, they found a greater dispersion between curves for mortars of different w/c prepared in a Hobart mixer based on ASTM C305 [50], with the lower w/c mortars showing earlier heat release and accelerated setting. According to this, the more viscous mortar receives more energy input during the mixing phase which might produce a better dispersion (deflocculation) of the cement particles and a greater rate of initial hydration as observed in Figure 6 [52]. The magnitude of this effect is negligible at later ages. In addition, a more concentrated pore solution in the lower  $w/c$  system (M1) compared to M8 may also increase the initial rate of hydration of the mixture.

The mixture with inert filler (M5) has 14 % more heat release at 7 d than that of M1. This percentage is the combination of the above mentioned 8 % (increased cement particle spacing) and an additional 6 % corresponding to the increased number of available nucleation sites and any extra cement surfaces exposed due to the deflocculation (assuming that fly ash and the inert filler have the same surface area and the same ability for providing nucleation sites/deflocculation). A faster rate of heat release and larger heat flow peak in M5 compared to M1 can also be observed in Figure 6, directly attributed to the provision of additional nucleation sites/exposed surface area.

When comparing M1 and M2 in Figure 5, the difference in heat release is 28 % at 7 d. This would correspond to increases of 14 % due to fly ash reaction, 6 % due to the increased number of nucleation sites/deflocculation, and 8 % due to the increased cement particle spacing. In addition to this, a 4-h retardation period is observed in M2 (Figure 6). It seems that this retardation effect is larger than the possible filler effect provided by the fly ash (as the first peak or main silicates reaction happens later than in M1). In this case, the fly ash likely reacts with some of the calcium ions in pore solution, diluting their concentration and thus lengthening the induction period [33], although retardation of early-age hydration by fly ash can also be due to other reasons, including sulfate imbalance. At the time that the required calcium concentration is achieved, the main silicates reaction takes place. Even though M5 and M2 have the same cement volume replacement level, M5 has a higher surface area than M2 (due to grinding of the inert filler). The higher surface area might provide a higher nucleation effect, accelerating the reaction. Also, the inert filler used in M5 has no absorption of water compared to the fly ash in M2 which might have a little absorption. This effect would leave more available water (liquid-filled space) in the M5 system than in M2, so that its cement particles might react faster.

In Figure 7, it can be observed that just the nucleation and spacing effects are detected during the first 45 h of hydration, the former starting to be measurable earlier than the latter. This can be explained as additional nucleation sites can provide templates for C-S-H precipitation before the increased spacing and water availability makes a measurable difference compared to M8. However, this temporal shift between both components of the physical effect of fly ash will depend on the particular characteristics of the system studied, such as  $w/c$  and replacement level. It is interesting to note that heat release caused by increased spacing (curve M8) increases over time, which can be explained by the fact that

the system with smaller spacing (M1) will have cement hydration products from different particles impinging on one another sooner and will thus release less and less heat than in a well-spaced system (M8). It is also observed that at about 45 h of hydration, the fly ash reactions start having a measurable effect. At 7 d, it can be observed that the physical (spacing, deflocculation, and nucleation) and chemical contributions of fly ash account for around 50 % each of the additional heat release of M2 compared to M1.

The same type of analysis can be done in the other two systems (60 % and 80 % volume replacement levels). Figures 8–11 show the results for these two systems. It can be noticed that the same trends as those in the 40 % system are observed in terms of increased heat release and acceleration/retardation effects, although with higher heat flow peaks and larger separation among heat curves as the volume replacement level increases. This is due to the type of normalization used in producing these plots (per g of cement).

Higher fly ash volume replacement levels (e.g., 60 % to 80 %) increase the sulfate demand, since fly ash provides more aluminates that can react with the limited quantity of sulfates (gypsum) present in the system. This can be observed in both Figures 9 and 11, where the spatial relationship between the gypsum depletion peak (typically the  $2<sup>nd</sup>$  peak or a shoulder on the  $1<sup>st</sup>$  peak) and the main silicate peak is modified relative to the systems without fly ash [53]. As such, the hydration reaction of these systems might not be optimum, and the heat release and heat flow might be influenced by sulfate depletion. The effect of adding more sulfates to the mixture with 60 % fly ash replacement (M3) on the heat release and heat flow can be observed in Figures 12 and 13, respectively. Sulfates were added as 50 % calcium sulfate dihydrate (i.e., gypsum) and 50 % bassanite (i.e., calcium sulfate hemihydrate or plaster), as described by Niemuth [53]. While a larger amount of heat release is observed (Figure 12), the main silicates hydration peak occurs at the same time as in the mixture with a lower sulfate content, thus not affecting its rate of reaction (Figure 13). As can be seen in the Figures, this particular fly ash causes an increasing retardation of the early-age hydration reactions as its volumetric replacement level is increased and this retardation is not corrected by a further addition of sulfate [21].

#### **7. CONCLUSIONS**

When fly ash is used in concrete it is generally believed that the fly ash will react (hydraulically or pozzolanically), providing increased reaction products, reduced porosity, and increased durability. In addition, fly ash can act as a filler, especially at very early ages, when it has not yet reacted chemically. The filler effect results in three factors that should be considered (in addition to potential dilution of the pore solution): 1) larger separation among cement particles, which promotes more complete cement hydration since more space is available, 2) partial deflocculation of the cement particle 3-D network, providing increased exposed cement particle surface area, and 3) provision of new nucleation sites (on the filler) that can promote the formation of more hydration products.

This paper has attempted to infer, from isothermal calorimetry measurements, the physical and chemical effects that high volumes of fly ash have on the cement hydration. This was done in high volume fly ash and chemically inert filler mortar mixtures. Three cement

volume replacement levels have been used for this purpose: 40 %, 60 %, and 80 %. The cement was replaced with either fly ash or an inert filler (silica sand) with a similar gradation as that of the fly ash. By using the chemically inert filler, the effects of increased cement particle spacing and additional nucleation sites were isolated from the chemical effect of fly ash reactivity.

The results indicate that the increased spacing of cement and nucleation/deflocculation effects provided by the addition of fly ash are dominant when compared with the chemical effect (i.e., fly ash reactivity) at early ages (less than 48 h), for the mixtures with a 40 % volume replacement level. This effect is even greater in mixtures with higher fly ash replacement levels. At early ages, the addition of the fly ash chosen for this study caused retardation in the system, as observed in the measured heat flow. The retardation is larger as the fly ash replacement level increases. After 48 h, the mixtures containing fly ash exhibited a greater cumulative heat release, which indicates a greater extent of hydration/reaction.

The approach shown in this study decouples the physical and chemical effects of fly ash.

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Examples of measured heat release in cement and cement-fly ash blended pastes. Uncertainty for these values will be presented later in the text (Figure 5 discussion).



#### **Figure 2.**

(a)  $w/c = 0.30$  plain system, (b) 60 % fly ash (or inert filler) system, and (c)  $w/c = 0.67$  plain system.





Cumulative PSD of the fly ash, inert filler blend, and the two inert filler fractions combined to form the inert filler blend.



#### **Figure 5.**

Heat release as a function of time. Comparison between  $w/c = 0.3$  plain,  $w/cm = 0.3 + 40$  % fly ash or inert filler, and  $w/c = 0.47$  plain mortar mixtures.



**Figure 6.** 

Heat flow as a function of time. Comparison between  $w/c = 0.3$  plain,  $w/cm = 0.3 + 40$  % fly ash or inert filler, and  $w/c = 0.47$  plain mortar mixtures.

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#### **Figure 7.**

Additional heat (in reference to M1) attributed to increased spacing factor, additional nucleation sites, and fly ash reaction in a  $w/cm = 0.3$  mortar with 40 % of the cement volume being replaced.



#### **Figure 8.**

Heat release as a function of time. Comparison between  $w/c = 0.3$  plain,  $w/cm = 0.3 + 60$  % fly ash or inert filler, and  $w/c = 0.67$  plain mortar mixtures.

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**Figure 9.** 

Heat flow as a function of time. Comparison between  $w/c = 0.3$  plain,  $w/cm = 0.3 + 60$ % fly ash or inert filler, and  $w/c = 0.67$  plain mortar mixtures.



#### **Figure 10.**

Heat release as a function of time. Comparison between  $w/c = 0.3$  plain, and  $w/cm = 0.3 +$ 80 % fly ash or inert filler.

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Heat flow as a function of time. Comparison between  $w/c = 0.3$  plain, and  $w/cm = 0.3 + 80$ % fly ash or inert filler.

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**Figure 12.**  Effect of sulfate addition on the heat release of a 60 % fly ash mortar mixture.

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**Figure 13.**  Effect of sulfate addition on the heat flow of a 60 % fly ash mortar mixture.

#### **Table 1.**

Chemical composition (wt. %) of the cement, fly ash, fine aggregate, and inert filler. (Uncertainty was determined to be  $\pm$  0.1 for values  $\pm$  0.1 %, and  $\pm$  0.01 for values < 0.1 %)



#### **Table 2.**

Mortar mixture proportions used in the study



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