Abstract: This paper reports the effect of addition of Polypropylene (PP) fibers in the performance of Self-Compacting Concrete (SCC) under elevated temperatures. Various SCC mixtures were made with PP fiber at 0.0, 0.05, 0.10 and 0.15% by volume added. The specimens were exposed to elevated temperatures (200°C [392°F], 400°C [752°F] and 600°C [1112°F]). The microstructure of the matrix was investigated. This study concluded that the high temperatures have a coarsening effect on the microstructure of SCC with the addition of PP fibers. It was reported that PP fibers reduced explosive spalling of SCC above 0.05% by volume and increased the number of voids in the microstructure significantly.

Keywords: Self-Compacting Concrete, Microstructure Coarsening, Elevated Temperatures, Polypropylene Fiber

Introduction

The process of investigating and collecting data related to the materials, products, structures, or components that failed is vital to the field of forensic engineering. This engrosses inspections, collecting evidence, measurements, developing models, obtaining exemplar products and performing experiments.

Fire in buildings and structures consumes lives and properties. Normal concrete is used worldwide as the main material in structures and buildings, which has led to a greater need to fully understand the effects of fire on concrete. Many researchers have investigated and documented the behavior of concrete under fire and concrete was found to have good fire resistance (Metey and Veiko, 1998). However, the behavior of SCC and HPC in fire has not yet been fully understood (Mehta and Monteiro, 1993).

The behavior of concrete at elevated temperature depends on its constituent materials, heating rate and peak temperature. Under certain heating conditions, the dehydration of C-S-H gel, the thermal incompatibility between the aggregate, cement paste and pore pressure within the cement paste are the main detrimental factors (Jahren, 1989). HPC is prone to spalling under high temperatures (Mehta and Monteiro, 1993). Concrete is found to lose its mechanical strength when subjected to severe spalling and cracking at elevated temperatures (Lea, 1920; Neves et al., 1997). Furthermore, exposure of concrete to high temperatures may also cause a change in pore structure, i.e., pore-structure coarsening (Al-Mutairi and Al-Shaleh, 1997).

Crook and Murray (1970) presented an experimental study on the behavior of conventional vibrated high strength concrete and self-compacting high strength concrete at high temperature with and without PP fiber. They used cylindrical and prismatic specimens at a heating rate of 0.5°C/min (32.9°F/min) (up to 400°C [752°F]) and high heating of up to 600°C (1112°F). They concluded that the residual mechanical properties of Self-compacting High Strength Concretes (SHSC) were similar to that of Conventional High Strength Concrete (CHSC). Spalling risk of SHSC was greater than for CHSC. The use of Polypropylene (PP) fiber improved the thermal stability of CHSC and SHSC. The addition of PP fiber modified the thermal gradient in tested concrete specimens during heating-cooling cycles.

The effect of temperature on different types of High-Strength Concrete (HSC) properties was presented (Kanéma et al., 2011). Thermal properties (specific heat, thermal conductivity and thermal expansion) are measured for these concrete types, namely, HSC, Self-Consolidating Concrete (SCC) and Fly Ash Concrete (FAC), at a temperature range from
20°C (68°F)–800°C (1472°F). The fibers’ (steel, polypropylene and hybrid) effect on thermal properties of HSC and SCC is also investigated. Results from their experiments showed that SCC possesses higher thermal conductivity, specific heat and thermal expansion than HSC and FAC in the 20°C (68°F)–800°C (1472°F) temperature range. Noumowé et al. (2006) studied the explosive spalling of concrete at elevated temperature subjected to heating-cooling cycles from room temperature to 600°C (1112°F). Two levels of heating rates (i.e., 0.1 and 1°C/min) with a fixed time of 1 h at the maximum temperature and a free cooling were imposed. The results showed that there was a link between the evolution of the thermal gradient in the specimens and the specimens’ weight losses. Their results also showed that the thermal stability of the concrete mixes was a function of the w/c ratio, heating rate and specimen dimensions. The factors causing explosive spalling are low permeability, weak water departure from the concrete and an increase of the thermal gradient. They also concluded that the thermal gradient (i.e., thermal stress) alone could not explain the explosive spalling of concrete.

The effects of elevated temperatures up to 800°C (1472°F) on compressive strength at different water-cement ratios of SCC were studied (Kanéma et al., 2011). They found that the hot compressive strength of SCC decreases with increasing temperature. Compared with normal-strength SCC, high-strength SCC possessed a larger compressive strength when exposed to high temperature. In addition, they found that the addition of PP fibers decreased the strength and probability of explosive spalling. Peng and Huang (2008) studied the change in microstructure of hardened cement paste (hcp) at elevated temperatures using X-ray diffraction, mercury intrusion porosimetry tests and Scanning Electron Microscope (SEM) observations. The decomposition of C-S-H at 600°C (1112°F), 700°C (1992°F) and 800°C (1472°F) were proposed. The C-S-H decomposition started at 560°C with significant effect above 600°C (1112°F). This indicated that the loss of strength by concrete exposed to temperatures above 600°C (1112°F) was related to the decomposition of HCP. They concluded that the pore size distributions confirm the coarsening effect of elevated temperatures on the pore structure, which was confirmed by the SEM analysis. The coarsening of pore structure can be regarded as the formation of equivalent cracks, which should be responsible for the reducing the HPC mechanical properties.

Nazari and Riahi (2010) investigated the microstructure, thermal properties and flexural strength of SCC with different amounts of SiO2 nanoparticles. They found that a partial replacement of cement up to 4 wt% SiO nanoparticle could accelerate C-S-H gel formation and increase crystalline Ca(OH)2 amount at the early ages of hydration. They concluded that SiO2 nanoparticles could shift the distributed pores to harmless and improve the pore structure of concrete.

Da Silva and de Brito (2015) characterized fresh state by a decrease of the coarse aggregates and a higher volume of mortar. An experimental programme was conducted to assess the effect of incorporating Fly Ash (FA) and Limestone Filler (LF) on the concrete’s porosity and the microstructure in binary and ternary mixes of SCC. The results enabled conclusions to be established regarding the SCC’s durability, based on its permeability and the microstructure of its pore structure.

The main objective of this research is to investigate the microstructure of SCC with PP fibers to mitigate spalling of concrete at elevated temperatures by using scanning electron microscope. The significance of this research is to study the microstructure of concrete at elevated temperatures using a scanning electron microscope.

Experimental Investigation

Materials and Mixture Proportion

Table 1 shows plain SCC mix was designed with constituent materials. PP fiber was then added to the control mix with 0, 0.05, 0.10 and 0.15% by volume of mix. Ordinary Portland Cement was used in the investigation. The specific gravity of cement was 3.15 and Blaine’s fineness was 2910 cm² g⁻¹. Class F fly ash from the Kapar Thermal Power Station, Malaysia, was used as a filler. The fly ash had a specific gravity of 2.32 and Blaine’s fineness of 2423 cm² g⁻¹ (28041.84 ft²/oz). Crushed angular granite of 20 mm (0.0656ft) maximum size was used as coarse aggregate. The specific gravity was 2.6, fineness modulus was 6.05 and a bulk density of 1480 kg/m³ (92.40lb/ft³) which conforms to an ASTM C 33-86 was used. The fine aggregates consisted of stream sand with a maximum size of 4.75 mm (0.1558ft), with normal grading, with a fineness modulus of 2-3. The absorption was 6.0% and the specific gravity was 2.5. Short PP fibers of length 19mm were used in the investigation with a density of 0.91 g/cm³ (1.244oz/ft³), a melting temperature of 160°C (320°F), a vaporization temperature of 341°C (645.8°F) and a burning temperature of 460°Cs (860°F) given by the supplier. Poly Carboxylic Ether (PCE) based superplasticizer, which is brown in color and a free flowing liquid and has a specific gravity of 1.15 superplasticizer, according to ASTM C 494-92. Type A and Type F were in aqueous form to water retention and to enhance workability. Potable tap water was used for mixing and curing.
Casting and Curing

All concrete mixes were prepared in 40L (1.41ft³) batches in a rotating planetary mixer. The batching sequence consisted of homogenizing the coarse aggregate and sand for 30 sec, then adding about half of the mixing water into the mixer and continuing to mix for 60 sec more. The mixer was then covered with plastic cover to reduce the evaporation of the mixing water and to let the dry coarse aggregates in the mixer absorb the water. After 5 min, the fly ash and cement were added and mixed for another 60 sec. Finally, the SP and the remaining water were introduced and the mixture was mixed for 3 min. Then, the fiber was added gradually and within 2 min PP fibers were added separately and with different percentages (0.0, 0.05, 0.10 and 0.15% by volume) to the target design mix to be tested under fire.

Samples of cubic (150×150×150 mm) (5.9×5.9×5.9 inch), cylinders (150×300 mm) (5.9×11.81 inch) and beams (100×100×500 mm) (3.94×3.94×5.9ft) specimens were casted and cured in water at 20°C (68°F) for 89 days to represent the behavior of wet condition on concrete. They were then tested at 90 days to determine the mechanical properties at temperatures within the normal range, 200°C (392°F), 400°C (752°F) and 600°C (1112°F).

Testing at Elevated Temperatures

The specimens were placed in an electrical box furnace of chamber size 600×300×300 mm (with a maximum temperature of 1200°C [2192°F]) and a heating rate of 5-10°C (50°F) per minute at elevated temperature. Concrete specimens were taken out from the curing environment at 89 days and kept under dry conditions in the laboratory for one day before being exposed to fire test. The temperature was kept at 200°C (392°F), 400°C (752°F) and 600°C (1112°F), each maintained for 4 h. The temperature was raised slowly until the required temperature was reached. The temperature was then maintained constant for 4 h. The specimens were then removed out of the oven, cooled down to room temperature by keeping them at room temperature, and exposing them to air circulation by a fan, then they were weighed and coded. Figure 1 shows the specimens in the furnace.

Table 1. Plain SCC mix with constituent materials

| Materials                     | Concrete mixture number |
|-------------------------------|-------------------------|
|                              | M0.0                   | M0.05                  | M0.10                   | M0.15                   |
| Cement, kg/m³ (lb/ft³)        | 437.5 (27.31)          | 437.5 (27.31)          | 437.5 (27.31)           | 437.5 (27.31)           |
| Fly Ash, kg/m³ (lb/ft³)       | 120 (7.49)             | 120 (7.49)             | 120 (7.49)              | 120 (7.49)              |
| Coarse Aggregate, kg/m³ (lb/ft³) | 730 (45.57)         | 730 (45.57)           | 730 (45.57)             | 730 (45.57)             |
| Fine Aggregate, kg/m³ (lb/ft³) | 907 (56.62)          | 907 (56.62)           | 907 (56.62)             | 907 (56.62)             |
| Water, kg/m³ (lb/ft³)         | 178 (11.11)            | 178 (11.11)           | 178 (11.11)             | 178 (11.11)             |
| w/p (ratio)                   | 0.32                   | 0.32                   | 0.32                    | 0.32                    |
| Superplasticizer, kg/m³ (lb/ft³) | 8.1 (0.506)          | 8.1 (0.506)           | 8.1 (0.506)             | 8.1 (0.506)             |
| Polypropylene fibers, % by volume of mix | 0.0                  | 0.05                  | 0.10                    | 0.15                    |

Fig. 1. The specimens in the furnace
Fig. 2. Scanning electron microscopy of plain SCC from (a) to (e), (a) Plain SCC at Room Temperature (27°C), (b) Plain SCC at 600°C, (c) Plain SCC at room temperature (27°C), (d) 4 h, 400°C and plain SCC, (e) 4 h, 600°C and plain SCC

The specimens are broken at 90 days to permit an evaluation of the mix. The real dimensions of the test specimens and the maximum applied load were measured. The micro structure of SCC mixtures was tested on a small specimen of concrete with a diameter of approximately 25 mm (1 inch) and a thickness of about 4 mm (0.16 inch), and the specimen had been cut in order to be tested by SEM. The specimens were previously coated with a very thin layer of gold to prevent charge built-up on the surface after being subjected to elevated temperature (Fig. 2).

Results and Discussion

Spalling Assessment

Figure 3 shows specimens subjected to fire test. In Fig. 3a, no spalling occurs although there were specimens of plain SCC, but there were small cracks on the surface of the specimens when the temperature was 200°C (392°F) for a heating period of 2 h.

Also in Fig. 3b, specimens of plain SCC showed more cracks appeared on the surface when the time of exposure was increased to 4 h. In Fig. 3c, plain SCC specimens showed small cracks appear and no explosive spalling occurred with peak temperature maintained at 400°C (752°F) for 4 h.

Figure 4 shows some of the specimens after heating with peak temperature maintained at 600°C (1112°F) for 2 and 4 h. In Fig. 4a, the most common type of damage was in specimen number 2, as nearly everything from the middle to the upper of the specimens were exploded; in specimen number 3, explosive spalling took place in the upper part as one big piece; and specimen number 1 did not spall. In other modes, Fig. 4b nearly behaved in the same way and showed the same trend because they were tested under the same temperature, which was 600°C (1112°F). All these tests were conducted on plain SCC specimens at 600°C (1112°F) with an exposure time of 2 and 4 h.
Fig. 3. Plain SCC mixtures specimens, (a) 2 h, 200°C and plain SCC, no explosive spalling, (b) 4 h, 200°C and plain SCC, no explosive spalling, (c) 4 h, 400°C and plain SCC, no explosive spalling
A compression with degree of gradation was done by Equation 1 (Da Silva and de Brito, 2015) to identify the criterion used to quantify the spalling severity:

\[
Spalling\ \text{degree} = \frac{m_1}{m_2}
\]

(1)

Where:

- \(m_1\) = Mass of concrete lost due to spalling
- \(m_2\) = Mass of the specimen before testing

If the spalling degree = 0.3, then 30% of the specimen mass was lost as a direct result of spalling. For the test on specimens with plain SCC and a 2 h exposure time and at 600°C (1112°F), the spalling was measured as in Equation 1 as follows:

\[
Spalling\ \text{degree} = \frac{0.24}{1.58} = 0.15
\]

This means that 15% of the cylinder specimens are lost as a direct result of spalling. Also, for a test on cylinder specimen at plain SCC fibers over a 4 h exposure time and at 600°C (1112°F), the spalling is measured as follows:

\[
Spalling\ \text{degree} = \frac{0.5}{1.6} = 0.31
\]

This means that 31% of the cylinder specimens are lost as a direct result of spalling.

Scanning Electron Microscopy of Different Percentages of Polypropylene Fibers

Figure 5a-h show the matrix SCC with PP fibers and granite aggregate adherence determined under scanning through electronic microscope for 0.0, 0.05, 0.10 and 0.15% of PP fibers by volume of SCC mixture.

The permeability and mineralogy of the aggregate may influence concrete when exposed to fire. Depending on the rate of heating, permeability, size and condition of the aggregate, the porous aggregate may be susceptible to disruptive expansion. Granite, which is formed from siliceous aggregate containing quartz, can distress in concrete at a temperature of 500°C (932°F) to 700°C (1292°F). The loss is due to decomposition of calcium hydroxide in the cement paste as well as the decomposition of calcium silicate hydrate compound (C-S-H) phases. This is followed by the formation of β-C2S (Schneider et al., 1982). The mineralogy of aggregate determines the differential thermal expansions between the cement paste, the aggregate and the ultimate strength of the interfacial transition zone.

The temperature effect on hydrated cement paste depends on the degree of hydration and the moisture state. It consists mainly of calcium sulphaaluminate hydrates, calcium silicate hydrate, and calcium hydroxide. A saturated paste contains great amounts of capillary and adsorbed water, so that when the temperature rises the water will vaporize to steam. The temperature of the concrete will not rise until all the evaporable water has been removed. SCC contains constituent materials with a small size that makes the cement paste impermeable such that it can conserve the water inside. With an increasing rate of temperature, damage may take place in the form of explosive spalling. This occurs when the vapor pressure of the steam inside the cement paste increases at a rate greater than the tensile strength of the cement paste.

Figure 5a to h indicates that at 2 and 4 h exposure times and with the increase in temperature to 200°C (392°F), 400°C (752°F), 600°C (1112°F) and at 200°C (392°F) to 300°C (572°F), the interlayer C-S-H water and some of the chemically combined water from the C-S-H and sulphaaluminate hydrates would also be lost. In addition, further dehydration of the cement paste due to decomposition of calcium hydroxide begins at about 500°C (932°F) to 600°C (1112°F).
Fig. 5. Scanning Electron Microscopy of SCC from (a) to (h), (a) 2 h, 200°C and 0.10% PP, (b) 2 h, 200°C and 0.15% PP, (c) 2 h, 400°C and 0.05% PP, (d) 2 h, 400°C and 0.10% PP, (e) 2 h, 600°C, 0.05% PP, (f) 2 h, 400°C and 0.15% PP, (g) 2 h, 600°C and 0.15% PP, (h) 4 h, 200°C and 0.05% PP
Spalling Assessment of Percentages of Polypropylene

Figure 6a and b shows that addition of PP fibers could be used to mitigate spalling. As seen in Fig. 6a, the SCC with 0.05% of PP fibers with peak temperature maintained at 400°C (752°F) for 4 h no spalling occurred. Furthermore, in Fig. 6b, as the peak temperature was maintained at 600°C (1112°F) for 4 h time of exposure, no explosive spalling occurred.

Scanning Electron Microscopy General Assessment

In general, it is concluded that a study using SEM was completed to reveal the reaction of the SCC to heated temperatures. Images of the SCC exposed to lower temperatures of 70°C (158°F) show that ettringite occurred. The loss of free water takes place at temperatures greater than 105°C (221°F). The PP melts at 162°C (323.6°F) and also vaporizes as the temperature and exposure time increases. When the test cylinders cool down for testing, the melted PP fibers become hardened and brittle, as they are allowed to remain in the cylinders to give them strength. Oxidation of the iron oxide in the aggregate occurs at about 300°C (752°F) and the dissociation of the sand particles in parts of the siliceous aggregate can be observed in the images. The dissociation of Portland-ite at 400°C (752°F) to 600°C (1112°F) can also be seen. At 600°C (1112°F) more PP fibers vaporize, leaving less liquid available in the molds and so the samples exhibit less strength at higher temperatures.

Also, changes were noted in the volume associated with water held by the small pores and the melting of the PP fibers at 162°C (323.6°F) with respect to the water vapor that comes from capillary water with voids larger than 0.05 µm and from absorbed water that is held close to the solid surface of hydrated cement paste by attractive forces; water integrated into C-S-H structure and water in the microstructure-these all evaporate, then the voids will increase as testing temperature increases from 27°C (80.6°F), 200°C (392°F), 400°C (752°F), to 600°C (1112°F). The melting and releasing of PP fibers to the surface of the sample can be seen in the images as they form channelization for the water vapor to flow outside the sample. This phenomenon mitigates explosive spalling that can occur in high performance concrete, of which SCC is an example.

Conclusion

The following conclusions can be derived from the investigation on the behavior of SCC at elevated temperature.

SEM can reveal the reaction of the samples when heated. Images of samples.

Exposure to low temperatures below 70°C (158°F) show that ettringite occurs. The loss of free water takes place at temperatures greater than 105°C (221°F). The PP melts at 162°C (323.6°F) and also vaporizes as the temperature and exposure time increases. When the test cylinders cool down for testing, the melted PP fibers become hardened and brittle, as they are allowed to remain in the cylinders to give them strength. Oxidation of the iron oxide in the aggregate occurs at about 300°C (752°F) and the dissociation of the sand particles in parts of the siliceous aggregate can be observed in the images. The dissociation of Portland-ite at 400°C (752°F) to 600°C (1112°F) can also be seen. At 600°C (1112°F) more PP fibers vaporize, leaving less liquid available in the molds and so the samples exhibit less strength at higher temperatures.

The changes in volume associated with water held by small pores and the melting of PP fibers at 162°C (323.6°F) with respect to the water vapor that comes from capillary water with voids larger than 0.05 µm and from absorbed water that is held close to the solid surface of hydrated cement paste by attractive forces; water integrated into C-S-H structure and water in the microstructure-these all evaporate, then the voids will increase as testing temperature increases from 27°C (80.6°F), 200°C (392°F), 400°C (752°F), to 600°C (1112°F). The melting and releasing of PP fibers to the surface of the sample was seen in the images to form channelization for water vapor to flow outside the sample. This phenomenon mitigates explosive spalling that occurs in high performance concrete and SCC is one of them as seen from the images of the microscopy tests.
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Author’s Contributions

Mahmoud B. Alhasanat and Arabi N. Al Qadi: Participated in all experiments, coordinated the data-analysis and contributed to the writing of the manuscript.

Omar A. Al Khashman and Ahmad Dahamsheh: Participated in all experiments, coordinated the data-analysis.

Ethics

This article is original and contains unpublished material. The corresponding author confirms that all of the other authors have read and approved the manuscript and no ethical issues involved.

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