SHRINKAGE REDUCTION AND CRACK PREVENTION OF ALKALI-ACTIVATED PHOSPHOROUS SLAG CEMENT

Yanan Wang,* Yonghao Fang, Yongfan Gong, Chenhui Zhu, Yamin Gu

College of Mechanics and Materials, Hohai University, Nanjing 210098, P.R. China

*E-mail: fangyh@hhu.edu.cn

Submitted September 20, 2015; accepted February 1, 2016

Keywords: Alkali-activated cement, Phosphorous slag, Shrinkage, Cracking, Expansive agent, Polypropylene fiber

The effects of fly ash, calcium oxide and polypropylene fiber on the physical and mechanical properties, shrinkage and cracking behaviors of alkali-activated phosphorous slag cement (AA–PS–C) were studied. The results show that replacing 10 - 15 % phosphorous slag by fly ash and adding calcium oxide as an expansive agent reduce the shrinkage of AA–PS–C. Fly ash will increase the flexural strength, although the compressive strength will be slightly decreased, while the calcium oxide expansive agent coated with aluminum stearate will slightly shorten the setting time and reduce the strength. Adding polypropylene fiber can greatly increase the crack-resistance of AA–PS–C.

INTRODUCTION

Phosphorous slag is a waste by-product from yellow phosphor production. Research results have proved that it is possible to prepare alkali-activated phosphorous slag cement (AA–PS–C) by mixing phosphorous slag powder, or mixture of that with fly ash, with water glass [1-3]. These cements exhibit high strength, and their corrosion and frost resistances are significantly better than those of Portland cement. Unfortunately, pastes prepared with AA–PS–C exhibit high shrinkage, much higher than pastes prepared with ordinary Portland cement. As a consequence, cracking of the cement paste and concrete occurs just as in alkali-activated blast furnace slag cement [4-8]. Although expansive agents and shrinkage-reducing admixtures have been efficiently used in ordinary Portland cement concrete [9-11], they are not effective for AA–PS–C. Either they are inefficient or have severe adverse effects on the mechanical properties [12].

To reduce the shrinkage of AA–PS–C, the effects of fly ash partly replacing phosphorous slag, and addition of calcium oxide expansive agent (CE) on the shrinkage, setting time and strength of AA–PS–C were studied in the present work. Polypropylene fiber has been widely and efficiently used in cracking prevention of ordinary Portland cement concrete [13-16]. Therefore, in the present work, the cracking prevention effect of polypropylene fiber on AA–PS–C was studied as well.

EXPERIMENTAL

Materials

Granulated phosphorous slag (PS) with Blaine specific surface area (SSA) of 483 m$^2$·kg$^{-1}$ was supplied by Yunnan Phosphorus Group Co., Ltd, and fly ash (FA) with SSA of 462 m$^2$·kg$^{-1}$ was supplied by Nanjing Huaneng Power Plant, respectively. The chemical compositions of PS and FA as analyzed by Jiangsu Test and Analysis Center with chemical methods according to JC/T 1088-2008 [17] and GB/T 176-2008 [18], respectively, are shown in Table 1. The SiO$_2$/Na$_2$O module of the water glass used as the activator was 1.4. Calcium oxide expansive agent (CE) was prepared by firing analytical reagent CaCO$_3$ at 1450°C for 2 hours, followed by mixing with 0.5 wt. % aluminum stearate at 200°C, so that the aluminum stearate melted and coated the surface of the CaO particles. Ordinary Portland cement P·II 42.5 (PII), conforming to the Chinese standard GB 175-2007 [19], were used as the reference. ISO standard

<table>
<thead>
<tr>
<th>Material</th>
<th>SiO$_2$</th>
<th>Al$_2$O$_3$</th>
<th>Fe$_2$O$_3$</th>
<th>CaO</th>
<th>MgO</th>
<th>P$_2$O$_5$</th>
<th>SO$_3$</th>
<th>IL*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PS</td>
<td>41.23</td>
<td>4.40</td>
<td>4.87</td>
<td>44.06</td>
<td>0.27</td>
<td>1.44</td>
<td>–</td>
<td>1.70</td>
</tr>
<tr>
<td>FA</td>
<td>49.76</td>
<td>32.88</td>
<td>4.04</td>
<td>3.67</td>
<td>1.09</td>
<td>0.63</td>
<td>2.32</td>
<td></td>
</tr>
</tbody>
</table>

* Mass loss after being ignited at 950°C for 20 min.
sand was used for preparing mortar specimens. A type of polypropylene fiber with diameter and length 28 μm and 12 mm, respectively, was used to reduce cracking.

Experimental methods

Setting time and strength test

AA–PS–C cements with the raw material mix proportions shown in Table 2 were prepared. The content of water glass equivalent to Na$_2$O for all the cements was fixed at 6 wt. % of the sum of PS, FA and CE. The setting time of the cement pastes were tested according to GB/T 1346-2001 [20].

The flexural and compressive strengths were tested according to GB/T17671-1999 [21]. Mortar specimens of 40 × 40 × 160 mm were prepared with a sand to AA-PS-C ratio of 3.0 and a water (including the water contained in the water glass) to AA–PS–C ratio of 0.5. The specimens were cured in the mould at 20 (± 1)°C and relative humidity (RH) > 90 % for 24 hours, and cured in water of 20 (± 1)°C after demoulding. Flexural and compressive strengths were tested at the age of 3 days and 28 days by a TYE-10C flexural and compressive testing machine for cement mortar manufactured by China Wuxi Jianyi Instrument and Machinery Co., Ltd. Three specimens were tested each cement at each age according to the standard.

Table 2. Mix proportions (in weight) of AA–PS–C.

<table>
<thead>
<tr>
<th>Cement</th>
<th>PS</th>
<th>FA</th>
<th>CE</th>
</tr>
</thead>
<tbody>
<tr>
<td>PF0</td>
<td>100</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>PF1</td>
<td>90</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>PF2</td>
<td>85</td>
<td>15</td>
<td>0</td>
</tr>
<tr>
<td>PF3</td>
<td>80</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>PF4</td>
<td>70</td>
<td>30</td>
<td>0</td>
</tr>
<tr>
<td>PFC1</td>
<td>85</td>
<td>15</td>
<td>1</td>
</tr>
<tr>
<td>PFC2</td>
<td>85</td>
<td>15</td>
<td>2</td>
</tr>
<tr>
<td>PFC3</td>
<td>85</td>
<td>15</td>
<td>3</td>
</tr>
<tr>
<td>PFC4</td>
<td>85</td>
<td>15</td>
<td>4</td>
</tr>
</tbody>
</table>

Table 3. Physical and mechanical properties of AA–PS–C.

<table>
<thead>
<tr>
<th>No.</th>
<th>Water requirement (%)</th>
<th>Setting time (min)</th>
<th>Flexural strength (MPa)</th>
<th>Compressive strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>initial</td>
<td>final</td>
<td>3 day</td>
</tr>
<tr>
<td>PF0</td>
<td>26.4</td>
<td>50</td>
<td>75</td>
<td>4.7</td>
</tr>
<tr>
<td>PF1</td>
<td>26.0</td>
<td>55</td>
<td>78</td>
<td>4.0</td>
</tr>
<tr>
<td>PF2</td>
<td>25.8</td>
<td>58</td>
<td>80</td>
<td>4.4</td>
</tr>
<tr>
<td>PF3</td>
<td>25.6</td>
<td>62</td>
<td>85</td>
<td>3.8</td>
</tr>
<tr>
<td>PF4</td>
<td>25.4</td>
<td>72</td>
<td>94</td>
<td>3.1</td>
</tr>
<tr>
<td>PFC1</td>
<td>26.6</td>
<td>55</td>
<td>72</td>
<td>4.3</td>
</tr>
<tr>
<td>PFC2</td>
<td>26.8</td>
<td>48</td>
<td>64</td>
<td>4.5</td>
</tr>
<tr>
<td>PFC3</td>
<td>27.0</td>
<td>46</td>
<td>58</td>
<td>4.2</td>
</tr>
<tr>
<td>PFC4</td>
<td>27.4</td>
<td>32</td>
<td>45</td>
<td>4.1</td>
</tr>
<tr>
<td>PFC3F*</td>
<td>Prepared by adding 0.3 wt.% of polypropylene fiber (based on the sum of phosphorous slag and fly ash).</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

RESULT AND DISCUSSION

Setting time and mortar strength

Shown in Table 3 are the results from the normal water requirement, setting time and strength tests. When the replacing quantity of fly ash for phosphorous slag increased (PF0-PF4), the setting time of the AA–PS–C increased. Addition of calcium oxide expansive agent (PFC1-PFC4) increased the water requirement of the cement and the test was conducted at 20 (± 2)°C with RH = 55 (± 3) % and the wind speed 0.5 m$^2\cdot$s$^{-1}$. The widths of the cracks were measured by a reading microscope with a magnification of 20, combined with a digital camera.

Shrinkage test

The shrinkage of the cement mortar specimens were tested according to JC/T 603-2004. Three specimens of 25 × 25 × 280 mm were molded for each mixture and moisture cured for 1 day at 20 (± 1)°C and 95 (± 2) % relative humidity (RH) in a curing chamber. Then the specimens were demolded and cured in water at 20 (± 1)°C for 2 days. The original lengths were measured at the age of 3 days when the specimens were taken out of the water. The specimens were then continued to be cured in the same condition for 6 days, and the moisture curing shrinkages were measured after 3, 5 and 7 days, respectively. After that, the specimens were cured at 20 (± 2)°C and RH = 60 (± 3) % in a drying test chamber. The drying shrinkages from 1 day up to 49 days were measured.

Cracking prevention test

The cracking prevention effect of polypropylene fiber for AA–PS–C was tested according to the mortar plate method of CECS 38: 2004 [22]. The dosage of the fiber was 0.3 % of the weight of the cement, and the test was conducted at 20 (± 2)°C with RH = 55 (± 3) % and the wind speed 0.5 m$^2\cdot$s$^{-1}$. The widths of the cracks were measured by a reading microscope with a magnification of 20, combined with a digital camera.
added as the alkaline activator. Since the calcium oxide expansive agent used in the present work was “dead-burnt”, the hydration activity was relatively low. In addition, the particle surface of the expansive agent was coated with aluminum stearate, which blocked the contact of particles with water, and retarded the hydration of CaO, so that the setting accelerating effect of the calcium oxide expansive agent was greatly reduced. Alkali-activated cement with an initial setting time longer than 45 minutes was prepared when the dosage of the calcium oxide expansive agent was not higher than 3 % of the sum of phosphorous slag and fly ash. When the dosage was higher, the setting time of the cement decreased to be shorter than 45 minutes.

The compressive strengths of pure AA–PS–C are very high after both 3 days and 28 days, while the flexural strengths are relatively lower. When fly ash was added as a replacement of the phosphorous slag, the compressive strength was significantly reduced, and the flexural strength after 3 days was also somewhat reduced, while the flexural strength after 28 days was increased when the replacement did not exceed 15 %. Since the activity of fly ash is lower than that of phosphorous slag, replacing phosphous slag with fly ash decreased the cementitous hydration products formed after comparable times, which resulted in a decrease of compressive strength. On the other hand, the fine particles of quartz and mullite in the fly ash can play the role of a “micro-aggregate”, which can, in a certain extent, block the expansion of cracks when the specimen was subjected to flexural load, and thus increase the flexural strength.

As the dosage of calcium oxide expansive agent increased, the strength of the cement decreased, though the decrement was not very significant when the dosage was not higher than 3 %. Adding 0.3 % polypropylene fiber based on the sum of phosphorous slag and fly ash (PFC3F) slightly decreased the 3-day strengths, while having little effect on the 28-day strengths.

Shrinkage of mortar

Figure 1a illustrates the change of shrinkage of the AA–PS–C mortars with the replacement of phosphorous slag by fly ash at different ages. The result for PII cement mortar is also shown in the figure as a reference. During the first 7-day moisture curing, the PII cement mortar nearly did not shrink, while almost all the alkali-activated slag cement mortars shrunk in different extents. During the dry curing stage, all the specimens shrunk significantly, and the AA–PS–C mortars shrunk much more than the PII cement mortar. The greatest total shrinkage after 56 days was as high as 0.21 % for PF0 mortar, 70 % higher than the PII cement mortar. When phosphorous slag was partly replaced by fly ash, the shrinkage at each age was reduced, the total shrinkages after 56 days for PF2 and PF3 with the replacement of 15 and 20 %, respectively, were reduced to be about 0.17 %. The shrinkage reduction effect of fly ash is ascribed to the low reactivity of fly ash compared with phosphorous slag powder, and the restricting effect of the inactive mineral and glassy particles in the fly ash.

Figure 1b shows the change of shrinkage of the AA–PS–C mortar specimens with the dosage of calcium oxide expansive agent at different conditions and different ages. The results for the PII cement and the pure AA–PS–C (PF0) are also shown for reference. In the first 7-day moist curing, the specimens to which calcium oxide expansive agent was added not only did not shrink, but even expanded. The expansion of the cement was caused by the fact the aluminum stearate coating on the surface of the calcium oxide expansive agent particles gradually dissolved in the alkaline solution of the cement, so that the block between the water and the calcium oxide expansive agent particles was removed. The latter reacted with water and calcium hydroxide formed, resulting in an increase of the solid volume,
which not only compensated the chemical shrinkage of the cement, but even caused a expansion. As the dosage of the calcium oxide expansive agent increased, the expansion of the mortars in the moist curing stage increased, while the shrinkage in the dry curing stage remained almost unchanged. The total shrinkage of the PFC3 mortar specimen with 15 % fly ash replacement and 3 % calcium oxide expansive agent dosage after 56 days was 0.12 %, being reduced by 40 % compared with that of PF0 with neither fly ash replacement nor calcium oxide expansive agent addition. The result confirms the desirable shrinkage reducing effect of the combination of fly ash and calcium oxide expansive agent.

Cracking prevention effect of polypropylene fiber

The shrinkage of cement-based materials results in stress. Cracking will take place when the stress exceeds the tensile strength of the materials. Alkali-activated slag cement is characterized by high shrinkage, so that its shrinkage reduction and cracking prevention demands more attention. Adding fibers, especially together with expansive agent, has turned out to be efficient in preventing cracking of ordinary cement concrete.

Figures 2a, 2b and 2c show the micrographs of the specimen surfaces, showing the cracking states of the plate mortar specimens of PF2, PFC3 and PFC3F, respectively, after 48 hours wind drying. PFC3F was prepared by adding 0.3 wt. % polypropylene fiber to PFC3. There are quite a few cracks with lengths of 5-20 mm and widths of 0.2 - 0.8 mm on the surface of the PF2 specimen (Figure 2a). When calcium oxide expansive agent was added, the cracking was greatly reduced, and both the number and the size of the cracks of the PFC3 are much smaller than those of the PF2 specimen (Figure 2b). It is the formation of calcium hydroxide that partly compensated the chemical and drying shrinkage of the cementitious materials. When both calcium oxide expansive agent and polypropylene fiber were added (PFC3F), only very few small cracks were observed in the specimen, and no significant change took place after 72 hours wind drying. The result indicates that the crack resistance of the alkali-activated cement has been greatly improved. The cracking restriction effect of the polypropylene fiber on the alkali-activated slag cement is based on the same mechanism as that in ordinary Portland cement-based materials [13].

CONCLUSIONS

- Replacing 15 - 20 % phosphorous slag by fly ash can increase the flexural strength and reduce the shrinkage of AA–PS–C.
- The calcium oxide expansive agent coated with aluminum stearate will slightly shorten the setting time and reduce the strength of AA–PS–C, but can greatly reduce the early age shrinkage.
- The combination of calcium oxide expansive agent and polypropylene fiber can greatly improve the crack resistance of AA–PS–C.

Figure 2. Cracking behavior of the mortar specimens; a) PF0, b) PFC3, c) PFC3F.
Acknowledgements

The authors would like to gratefully appreciate financial support from the National Natural Science Foundation of China (Grant No. 51461135001 and 51139001), and the Provincial Foundation for Industry-University-Research Joint Innovation Cooperation(Grant No. BY2014002-04) of Jiangsu, China.

REFERENCES


