Mechanical Properties, Pozzolanic Activity and Volume Stability of Copper Slag-Filled Cementitious Materials

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Copper slag is a by-product waste during copper pyrometallurgy. The piling-stack of copper slag not only occupies a large quantity of plowland, but causes potential danger for local residents. In order to reduce the storage of copper slag, this study prepared concretes by replacing ordinary cement by copper slag. The mechanical properties, phase composition, micrographs, pozzolanic activity and volume stability of the compound cementitious materials were evaluated with fly ash as control. It was found that when the content of copper slag was 5% ~ 10%, the best mechanical properties of concretes were obtained. The phase composition of hydration products of copper slag and fly ash based cementitious materials was similar. The micrographs and pozzolanic activity showed that copper slag had stronger reactivity than fly ash. Copper slag based pastes had good volume stability. Copper slag presented a potential application in mortars and concretes.

Keywords: mechanical properties, pozzolanic activity, volume stability, copper slag, cementitious materials.

1. INTRODUCTION

Copper slag (CS) is a metallurgy waste residue generated during the matte smelting process of copper metal [1]. When 1 t of copper is manufactured, 2.2 t of CS is generated [2]. In China, more than 1.500,000 t of CS is produced every year. Globally in the year 2015, approximately 68.7 million tons of CS was generated from the world copper industry [3]. These waste copper slags pile up like a mountain, occupying a large amount of plowland. The environmental pollution produced by the large quantity of CS affects the development of local mine areas. How to utilize CS effectively and cleanly, and push green construction, is a major project deserved to study in depth.

Some researches about the application of CS in concretes have been developed. On one hand, CS is studied to substitute the aggregates in mortars and concretes. The literatures show that when CS substitutes the sand in concretes, these concretes have the comparable physical properties with plain concretes [4 – 6]. The concretes with CS as fine aggregate have the nearly same compressive and tensile strength with plain concretes. Under proper substitution, the strength of the concretes with CS surpasses that of plain concretes. Brindha D, et al, reported that CS is in favour of the improvement of concrete durability [7]. On the other hand, besides the substitution of fine aggregates in concretes, CS can be used as mineral admixture and replace some quantities of cements, to cut down the costs and protect environment [8, 9]. I Alp, et al, studied the massive application of CS as raw material in cements [10]. Moura W A, et al, explored the properties such as compressive strength and water-absorbing capacity, and analysed the feasibility of CS used as pozzolanic material in concretes [11]. Caliskan and Behnood showed that the compressive strength of CS coarse aggregate concretes was marginally higher than that of limestone aggregate concretes [12]. However, most of the current researches that the compressive strength of CS coarse aggregate concretes was marginally higher than that of limestone paid attention to the mechanical properties of CS on mortars and concretes. There was a hand of literatures on the mechanism and active intensity of CS used as mineral admixture in cementitious system. In this study, the mechanical properties (including compressive, tensile, flexural strength and stress-strain relation), chemical activity and volume stability of cementitious system incorporating CS are explored. The results may supply more theoretical basis for the application of CS in mortars and concretes.

2. EXPERIMENTAL

2.1 Materials

Raw materials used in the experiment included copper slag (Jiangxi Tongxing copper factory, China), 42.5 grade Ordinary Portland Cement (OPC, Conch Cement Co., China) and fly ash (FA, class F, obtained from a coal fired power station in Zhengzhou, China). Fly ash powders were collected from coal smoke with a bag dust collector and dried at 105 °C for 24 h to reach constant weight before use. CS powders were obtained after 60 min of ball-milling on a planetary mill (QM-2SP20, Nanjing Nanda Instrument Plant, China) with a speed of 300 r/min.

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Table 1. Chemical composition (wt.%) and fineness (cm²/g) of OPC and mineral admixtures

<table>
<thead>
<tr>
<th>Sample</th>
<th>SiO₂</th>
<th>Al₂O₃</th>
<th>Fe₂O₃</th>
<th>MgO</th>
<th>CaO</th>
<th>Na₂O</th>
<th>K₂O</th>
<th>TiO₂</th>
<th>P₂O₅</th>
<th>MnO</th>
<th>SO₃</th>
<th>Fineness</th>
</tr>
</thead>
<tbody>
<tr>
<td>OPC</td>
<td>20.87</td>
<td>4.14</td>
<td>5.12</td>
<td>4.08</td>
<td>62.66</td>
<td>0.11</td>
<td>0.28</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>2.32</td>
<td>3289</td>
</tr>
<tr>
<td>CS</td>
<td>36.66</td>
<td>3.84</td>
<td>47.12</td>
<td>5.33</td>
<td>5.57</td>
<td>0.64</td>
<td>0.37</td>
<td>0.28</td>
<td>0.11</td>
<td>0.09</td>
<td>-</td>
<td>2874</td>
</tr>
<tr>
<td>FA</td>
<td>49.03</td>
<td>20.12</td>
<td>7.9</td>
<td>3.35</td>
<td>8.72</td>
<td>1.46</td>
<td>2.29</td>
<td>0.8</td>
<td>0.48</td>
<td>0.1</td>
<td>-</td>
<td>2047</td>
</tr>
</tbody>
</table>

Table 2. Mix design of CS concretes

<table>
<thead>
<tr>
<th>Sample</th>
<th>OPC, kg/m³</th>
<th>CS, kg/m³</th>
<th>Fine aggregate, kg/m³</th>
<th>Coarse aggregate, kg/m³</th>
<th>Water, kg/m³</th>
<th>Water-binder ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>C0</td>
<td>317</td>
<td>0</td>
<td>764</td>
<td>1062</td>
<td>190</td>
<td>0.6</td>
</tr>
<tr>
<td>C5</td>
<td>301</td>
<td>16</td>
<td>764</td>
<td>1062</td>
<td>190</td>
<td>0.6</td>
</tr>
<tr>
<td>C10</td>
<td>285</td>
<td>32</td>
<td>764</td>
<td>1062</td>
<td>190</td>
<td>0.6</td>
</tr>
<tr>
<td>C15</td>
<td>269</td>
<td>47</td>
<td>764</td>
<td>1062</td>
<td>190</td>
<td>0.6</td>
</tr>
</tbody>
</table>

The fineness of raw FA and milled CS was 2047 and 2874 cm²/g, respectively. The main chemical composition and particle fineness of the three raw materials were shown in Table 1. Fe₂O₃ and SiO₂ were the main chemical ingredients of CS, determined by X-ray fluorescence (XRF) spectrometry method. The phase composition of milled CS and raw FA was determined by X-ray diffraction (XRD) analysis. The main mineralogical phases were quartz and mullite in FA, while magnetite (Fe₃O₄) and fayalite (Fe₂SiO₄) were predominant in CS, shown in Table 2.

![Fig. 1. XRD patterns of milled CS and raw FA](image)

In the concrete experiment for mechanical properties, the OPC cements were substituted by CS in weight with different replacements (5%, 10% and 15%), shown in the mix design of Table 2. The water to binder (OPC+CS) ratio of 0.6 was used for all concrete admixtures with the slump of around 160 mm, according to the Chinese Standard GB/T 8007-2012 [14]. Deionized water was used to eliminate impurities influence.

Table 3. Composition of compound cementitious system

<table>
<thead>
<tr>
<th>Sample</th>
<th>CH, t.%</th>
<th>CS, wt.%</th>
<th>FA, wt.%</th>
<th>Water-binder ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS-CH</td>
<td>25</td>
<td>75</td>
<td>0</td>
<td>0.4</td>
</tr>
<tr>
<td>FA-CH</td>
<td>25</td>
<td>0</td>
<td>75</td>
<td>0.4</td>
</tr>
</tbody>
</table>

2.2. Methods

After curing age of 7 and 28 days in a standard curing chamber, the concrete samples with size of 100 mm × 100 mm × 100 mm were used to test compressive strength using a hydraulic 2000 kN press machine with the loading speed of 2500 N/s, following the Chinese Standard GB 50081-2002 [15]. The dimensions of samples for splitting tensile strength and flexural strength were 100 mm × 100 mm × 100 mm and 100 mm × 100 mm × 400 mm, respectively. For each curing age, triplicates for mechanical strengths were conducted. The stress-strain behavior was tested on the prism specimens (100 mm × 100 mm × 300 mm) after curing of 28 days, by using a MTS electronic universal testing machine with a capacity of 3000 kN. The load was controlled by displacement, and the loading rate was fixed at 0.003 mm/s.

The samples (20 mm × 20 mm × 20 mm) were prepared according to the mix of Table 3, removed after shaping for 1 day in a curing box, and then cured hermetically to predetermined curing time. The samples were immersed in acetone (Sinopharm, China) to stop hydration. After broken, grind, and dried, the samples were tested for X-ray diffraction (XRD), thermogravimetry-differential thermogravimetry/ differential scanning calorimetry (TG-DTG/DSC) and scan electron morphology (SEM). XRD was carried out by an X-ray diffractometer (X’Pert PRO, PANalytical Co., the Netherlands), employing CuKα radiation (1.54 Å). The instrument voltage and current were set as 40 kV and 40 mA, respectively. The 2θ range was from 5° to 60° with 0.013° step size and 10 s step time. The thermal analysis was carried out on a Mettler Toledo 850 apparatus at ranges of 35–1000°C (alumina crucible), with the heating rate being 10°C/min in air atmosphere. SEM micrographs were taken with a field emission scanning electron microscope (Nova 430, FEI, the
3. RESULTS AND DISCUSSION

3.1. Mechanical strengths

Fig. 2 demonstrates the mechanical strengths of concretes with different CS contents after 7 and 28 days of curing. In Fig. 2 a, with the increasing contents of CS, the compressive strength of concretes after curing of 7 days decreased, from 34.3 MPa of C0 to 28.7 MPa of C15. The compressive strength after curing of 28 days increased from 40.4 MPa of C0 and peaked with 43.2 MPa of C5, then decreased to 33.3 MPa of C15. The variation trend of tensile and flexural strengths of concretes with different CS contents after 7 and 28 days of curing was similar with the compressive strength of 28 days. The tensile and flexural strengths increased from C0 and peaked at C5, then decreased to C15. The tensile strength of C5 and C10 was nearly the same, while the flexural strength of C5 was slightly more than that of C10. Mineral admixtures can improve the concrete compressive strength, because of admixture chemical and physical effect [17]. The fine CS improved the particle distribution of cementitious system, and then enhanced the concrete compressive strength. Meanwhile, the copper slags with chemical activity reacted with CH of cement hydration products, which also enhanced the concrete compressive strength. However, the second hydration rate of mineral admixtures was relatively lower and mainly generated in the late hydration. The over CS substitution decreased the cement content and adversely affected the concrete compressive strength.

After 7 days of curing, the decreased compressive strength with the increased CS contents was due to the limited physical effect and low chemical activity of CS. This effect and activity could not remedy the adverse impact of decreased cement content. Whereas, after 28 days of curing, the increased mechanical strengths of concretes with 5% – 10% CS might be owe to the enhanced chemical activity of CS. But the too much replacement of CS at C15 sample could not bring the cement superiority because of the too limited content of Portland cements.

Brittleness is a kind of property of rupture with little deformation under external force. Normally, concretes have weak brittleness. In concrete engineering application and tests, brittleness index, namely the division of compressive strength by tensile strength (or flexural strength), is used to characterize the brittleness of concretes [18]. Fig. 3 shows the brittleness index of concretes with different CS contents. R_c/t denotes the brittleness index of the division of compressive strength by tensile strength, whilst R_c/f denotes that of the division of compressive strength by flexural strength. R_c/t expressed the similar trend with R_c/f, that the brittleness of 7 days curing decreased with the increased content of CS, reached the lowest at 10% CS content and increased when 15% CS content. In the early curing age, the compressive strength progressed fast and the formed C-S-H (hydrated calcium silicate) increased with the increased CS contents, which made the strong roughness. For 15% CS content, the less C-S-H was formed because of the over replacement of cements by CS. After 28 days of curing, the R_c/t and R_c/f expressed the same trend that the brittleness increased peaked at 5% CS, and then decreased. In the late curing age, the compressive strength increased slowly; while the formed C-S-H gel transformed to crystallizes.

Fig. 3. Brittleness index of concretes with different copper slag contents

Fig. 2. Mechanical strengths of concretes with different CS contents: a – compressive strength; b – tensile strength; c – flexural strength
Because of the least C-S-H gel in 5% CS specimen, its brittleness was the strongest and its roughness was the weakest. Also, the $R_{\alpha}$ and $R_{cfl}$ both can denote the brittleness index, due to the similar trend.

Fig. 4 depicts the uniaxial compressive stress-strain curve of concretes with different copper slag contents after 28 days of curing. The stress-strain curve demonstrated a typical brittle damage, with a smaller strain and a sudden fracture. Also, it can be seen that C5 had a higher compressive strength and elastic module than C0, demonstrating that the smaller content of copper slag can improve the compressive strength and elastic module of concretes. This might be due to the filling effect of copper slags in concrete. When OPC is replaced by copper slag in the same weight of 5 wt.%, copper slag can fill in the pores between hydration products and aggregates, and increase the compressive strength and elastic module. Our result is consistent with the literature [19]. For the breaking point in C5 stress-stain curve, the C5 concrete has less C-S-H gel produced by the less OPC, resulting in the bad ductility. When the replacement of copper slag in OPC surpasses a certain value, the elastic module of concretes with copper slag will decline, compared to concrete prepared by pure cement. Zain attributed the lower elastic module to the retardation of cement hydration due to the presence of heavy metals in copper slag [20]. Benson considered that the very fine particles of the slag supplied a large amount of surface area per unit volume to be coated with cement, which might effectively have reduced the amount of cement available for binding the fine and coarse aggregates [21]. Besides these reasons, we thought that for C10 specimens, the elastic module is lower than that of C0, possibly due to the less C-S-H produced in the concretes.

C$_3$S (tricalcium silicate) and C$_2$S (dicalcium silicate) in OPC can generate a large quantity of C-S-H gel and Ca(OH)$_2$ crystal by hydration reaction. Although active SiO$_2$ in copper slag can produce C-S-H gel via second hydration reaction with Ca(OH)$_2$ crystal, and Fe$_2$O$_3$ can hydrate to produce Fe(OH)$_3$/Fe(OH)$_2$ gel (clarified in Section 3.2), the gel is comparatively less when copper slag replaces OPC in the same weight.

The stress-strain curve, graphed constitutive relation, plays an important role in studying the structure and force analysis of concretes. So it is needed to construct the constitutive models. The stress-strain curve can be expressed as a dimensionless coordinate:

$$x = \frac{\varepsilon}{\varepsilon_c}, \quad y = \frac{\sigma}{\sigma_c}. \quad (1)$$

Thereinto, $\varepsilon$– the stress, $\varepsilon_c$– the peak stress, $\sigma$– the strain, $\sigma_c$– the peak strain. Then a standard curve with point coordinate (1,1) is plotted. Guo et al. deeply studied the constitutive relation of concretes and came up with a Guo’s model [22]. In Guo’s model, the stress-strain curve can be fitted as a polynomial for the ascent stage and a rational fraction for the descent stage:

$$
\begin{cases}
  y = ax + (3 - 2a)x^2 + (a - 2)x^3 & x \leq 1 \\
  y = \frac{x}{b(x - 1)^2 + x} & x \geq 1
\end{cases}
\quad (2)
$$

In Eq. 2, a and b are constants to be determined. By using Eq. 2, the constitutive relation of concretes with different copper slag contents is resolved. Because of the only ascent stage of the stress- strain curve, only the parameter, a, is determined, as shown in Table 4.

**Table 4. The value of parameter a**

<table>
<thead>
<tr>
<th>sample</th>
<th>C0</th>
<th>C5</th>
<th>C10</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>1.147</td>
<td>1.023</td>
<td>1.413</td>
</tr>
</tbody>
</table>

The stress-strain curves and the fitted curves in light of Eq. 2 are compared in Fig. 5. It can be seen the fitted curves nearly coincided with the tested stress-strain curves. It demonstrates that the Guo’s model can be used to analyze the constitutive relation of concretes with different copper slag contents.

### 3.2. Phase composition

If there were a quantity of active SiO$_2$ and Al$_2$O$_3$ in the mineral admixtures, these active composites could react with Ca(OH)$_2$ to generate C-S-H gel.
Such mineral admixtures were considered to have pozzolanic activity (good chemical activity). To verify the pozzolanic activity of CS, CS-CH samples were prepared in accordance to Table 3 with FA-CH as a control sample.

Fig. 6 shows the XRD patterns of CS-CH and FA-CH samples cured after 28 days. Besides the existence of CH in the hydration products of CS-CH and FA-CH, C-S-H and carbonated CaCO₃ were also detected, which demonstrated that CS had a kind of pozzolanic activity similar to FA. C-S-H was produced from the reaction of active SiO₂ in CS or FA and Ca(OH)₂.

![Fig. 6. XRD patterns of hydration products from CS-CH and FA-CH](image)

The presence of CaCO₃ was because of the carbonation of Ca(OH)₂ in atmosphere. Moreover, the characteristic peak intensity of CH in CS-CH hydration products was weaker than that of FA-CH, demonstrating that more CH in CS-CH cementitious system was consumed, and that the pozzolanic activity of CS was stronger than FA. The characteristic peak intensity of C-S-H in CS-CH hydration products was stronger than that of FA-CH, which was due to the faster reaction rate in more active CS-CH system, and the more hydration products (C-S-H) after 28 days of curing. Besides C-S-H gel found in the hydration products of CS-CH system, Fe(OH)₃/Fe(OH)₂ gel was generated, due to the Fe₂O₃ hydration reaction. The XRD patterns illustrated the stronger pozzolanic activity of CS than FA.

The DSC-TG curves of CS-CH and FA-CH after 28 days of curing is presented in Fig. 7. In the DSC curves, there are three obvious endothermic valleys between 30 °C and 1000 °C. The first endothermic valley happened below 200 °C, demonstrating the loss of absorbed water, the dehydration of C-S-H and Fe(OH)₃/Fe(OH)₂ gel [23, 24]. The second endothermic valley in 400 – 500 °C was due to the decomposition of unhydrated Ca(OH)₂. The last endothermic valley took place at about 700 °C, which meant the decomposition of CaCO₃. The existence of C-S-H gel, Ca(OH)₂ and CaCO₃ was in accordance to the XRD detection in Fig. 6. The area of the endothermic valley below 200 °C of CS-CH was bigger than that of FA-CH, which demonstrated more C-S-H gel in CS-CH than FA-CH. The endothermic valley in 400 – 500 °C in CS-CH was smaller than that of FA-CH, meaning that CS reacted with more Ca(OH)₂ and showed a stronger pozzolanic activity.

In DSC curves, the characteristic peaks in 400 – 500 °C were sole and clear. So the content, W₁, of Ca(OH)₂ could be calculated according to TG curves.

![Fig. 7. TG, DTG and DSC curves after 28 days of curing of: a – CS-CH; b – FA-CH](image)

In Table 3, the content of Ca(OH)₂ in the system including CS or FA, CH, and water was 17.9 wt.% (≈ 25 ÷ (100 ÷ 40) × 100 wt.%). Thus, the content of reacted Ca(OH)₂, α, was shown in Table 5. The bigger value of α meant the more intense hydration action in samples. After 28 days of curing, the reacted CH content (45.9 %) in CS-CH samples was more than the reacted CH content (13.0 %) in FA-CH samples, demonstrating the stronger pozzolanic activity of CS than FA.

**Table 5. Hydration degree of CS-CH and FA-CH**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Age (day)</th>
<th>W₁, %</th>
<th>α, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS-CH</td>
<td>28</td>
<td>9.7</td>
<td>45.9</td>
</tr>
<tr>
<td>FA-CH</td>
<td>28</td>
<td>15.5</td>
<td>13.0</td>
</tr>
</tbody>
</table>

### 3.3. Morphology of hydration products

Fig. 8 shows the SEM images of raw materials and hydration products after 28 days of curing. The CS presented irregular shape, with compact structure and rough surface, as shown in Fig. 8a. FA particles were spherical shape with glossy surface, as shown in Fig. 8c. Fig. 8b showed that after 28 days of curing, most surfaces of CS were obviously eroded, and compact gel substances were...
shaped on the CS surface, demonstrating that CS reacted with CH to generate C-S-H gel and enhanced the concrete compressive strength. However, as shown in Fig. 8 d, the FA particles were dispersed, and kept the original appearance, without being eroded, which illustrated that FA had weaker pozzolanic activity than CS.

Table 6. Volume stability of pastes with cement replaced by CS and FA at different contents

<table>
<thead>
<tr>
<th>CS content, wt.%</th>
<th>FA content, wt.%</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>20</td>
<td>50</td>
</tr>
<tr>
<td>30</td>
<td>50</td>
</tr>
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<td>40</td>
<td>50</td>
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<tr>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>60</td>
<td>50</td>
</tr>
</tbody>
</table>

Fig. 8. SEM images of raw materials and hydration products after 28 days of curing: a– raw material of CS; b– hydration products of CS-CH; c– raw material of FA; d– hydration products of FA-CH

3.4. Volume stability

In cementitious system, the content of CaO, MgO and limestone is much crucial to the volume stability of the system [25 – 27]. It could be seen from Table 1 that MgO in CS was 5.33 wt.%, with 3.35 wt.% in FA and 4.08 wt.% in cement. Therefore, it should consider the effect of MgO content on the volume stability of the cementitious system in the application of CS.

To explore the volume stability of CS cementitious system, cement was equally replaced by CS of 20 – 60 wt.%, or FA of 50 – 60 wt.%. The result was shown in Table 6. In plain cement, the instability of cement paste took place because of the volume expansion caused by Ca(OH)2 and Mg(OH)2, when f-CaO and f-MgO hydrated with water. Compared to plain cement, cement with mineral admixtures (CS or FA) had lower expansion rate (no more than 0.5%), which would not cause any instability. Some literatures reported that no f-CaO and f-MgO was found in CS. Nevertheless RO phase including divalent metal oxides, such as FeO, MgO, MnO and CaO, existed as solid solution in CS, which was more stable and would not hydrate even under high temperature and pressure [28, 29]. According to the aforementioned hydration tests, CS had strong pozzolanic activity, and could react with about half of Ca(OH)2 in the cement paste, which vastly reduced the Ca(OH)2 content. With the increase of CS in cement, the less of f-CaO and f-MgO, and the lower expansion rate of the cement paste. In a word, the cement paste with CS had good volume stability.

4. CONCLUSIONS

The feasibility of cement equally replacement by copper slag was explored on the mechanical strengths, pozzolanic activity and volume stability. The result showed that copper slag had stronger pozzolanic activity than fly ash and could react with Ca(OH)2 to generate C-S-H gel and enhance the concrete mechanical strength, when copper slag content was less than 15 wt.%. Compared to fly ash, the pozzolanic activity of copper slag was manifested earlier, and the reactive rate was faster. Moreover, the pozzolanic activity of copper slag was stronger, and copper slag could react with about half of Ca(OH)2 in the pastes. Though the high MgO content in copper slag, the cement pastes with copper slag had good volume stability, due to the lower Ca(OH)2 content in the pastes. These results manifested that copper slag had good mechanical strength, pozzolanic activity and volume stability, demonstrating that copper slag had potential application in cements and concretes.

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