ARTIFICIAL AGGREGATE FROM SINTERED COAL ASH

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Fly ash is one of the most commonly used secondary raw materials in the Czech Republic. It is used predominantly for re-cultivation, roads and additions to cement or plasters. The use of fly ash in the technology of sintering-based artificial aggregate was tested in the 1980s. However, the production was stopped for various technological and economic reasons. Nowadays, possibilities for the production of artificial aggregate are tested with fly ash produced in the Czech Republic and could be promising for future technologies, mainly with respect to the stability of production. The paper presents part of the study describing the possibilities of using microsilica and Fe2O3 for the optimization of a raw-material mix and fly-ash body. Three samples of high-temperature lignite combustion fly ash were selected from prospective sources in the Czech Republic. The fly ash was mixed with 5 % or 10 % additions. Then, samples were fired at temperatures of 1150 °C and 1200 °C. After firing, the physico-mechanical properties of the fly-ash bodies and microstructure were evaluated. The results imply that the addition of microsilica unambiguously improves the quality of the fly-ash body. The addition of Fe2O3 did not take part in the formation of the melt and weakened the fly-ash body’s structure.

Keywords: fly ash, ash body, firing, microsilica, sintering, iron trioxide

1 INTRODUCTION

Power plants producing electrical energy create energetic by-products during the combustion of pulverized lignite. The dominant proportion of the by-products is fly ash. Ecological and economic reasons motivate technological innovations focusing on using solid waste. The production of artificial fly-ash aggregate is a suitable construction material, which could be made from fly ash. Artificial, sintered, fly-ash aggregate is one of the few construction materials that can be produced only with fly ash. European and worldwide trends of newly developed building technologies accentuate the demand for the production of high-quality, sintered, artificial, fly-ash aggregate. If the character of the fly ash is optimal, no further treatment is needed. However, not every sample of fly ash has an optimal composition. The quality of fly ash has an influence on the composition of the mix, the technological parameters and the quality of the produced aggregate. For enhancing the properties of sintered fly-ash aggregate, additions commonly used in the ceramics industry could be used. The main candidates are microsilica and oxides of iron.

Microsilica is an amorphous SiO2, which is commonly obtained from separators during the production of ferrosilicon and silicon in electric arc furnaces. The application of microsilica in fire-resistant materials has been known for more than 40 years. The main task of microsilica in fire-resistant ceramic materials is the reaction in the system of binders, including the reaction mechanism at various temperatures. Various temperatures can be critical as regards the reactions. Microsilica consists of spheroidal particles with a mean diameter of around 0.15 microns. These spheroidal particles are a construction unit of primary agglomerates, which are bound to one another by strong bonds. The large specific surface and the wide distribution of microsilica increases the effectiveness of the encapsulation of the grains and the functionality of the fire-resistant ceramic materials compared to a narrow fraction. Microsilica is usually the finest part of the system with a specific surface of around 20 m² g⁻¹. The surface properties and possible impurities are important for a determination of the properties of the
final product. Microsilica can add more than 50 % to the total surface area of the particles in the mix.

Oxides of iron influence the formation of a ceramic body in various atmospheres, temperatures and firing agents for the production of sintered fly-ash aggregate. After granulation, the bodies are fired at a maximum temperature of around 1200 °C. After sufficient firing and cooling, the mix is crushed and screened for the final fractions.

For the production of aggregate of good quality, the granulometry of the input materials, their structure and chemical composition are also important. The produced sintered fly-ash aggregate is used for effective melting agents for the production of sintered fly-ash aggregate.

The optimal mix of appropriate raw materials for the production of artificial aggregate is mixed with water and granulated on a cylindrical or plate granulator, which makes the appropriate shape of the sintered fly-ash aggregate. After granulation, the bodies are fired at a maximum temperature of around 1200 °C. After sufficient firing and cooling, the mix is crushed and screened for the final fractions.

2 EXPERIMENTAL PART

Three samples representing prospective sources in the Czech Republic were selected for verification of the appropriateness of high-temperature lignite combustion fly ash for the production of artificial aggregate.

From the physico-mechanical and physico-chemical parameters we selected the loss on ignition (CSN 72 0103) showing unburned residues, the bulk density (CSN 72 2071), the specific surface area and the rest on the 0.045-mm sieve (CSN 72 2072-6). We also conducted the chemical and mineralogical analyses.

The parameters listed in Table 1 imply that fly ash produced in the Czech Republic has a minimal unburnt content and fulfills the requirements of the standard (CSN 72 2072-6, 2013) for a maximum loss on ignition of 15 %. For self-firing, it will be necessary to mix the fly ash with pulverized coal to achieve an optimal value of 8 % combustibles by weight. The greater coarseness of the fly ash FA3 evaluated with respect to the rest on the 0.0045-mm sieve and specific surface is evident. This fly ash also has a larger content of iron and lime, which can cause a reduction of the melting temperature of a batch. All the values of the high-temperature fly-ash samples fulfill the requirements of the Standard CSN 72 2072-6, 2013 for a minimal value of 800 kg m⁻³.

As is clear from Table 2 FA1 has the highest percentage of mullite and a lower content of amorphous phase in comparison with FA2 and FA3. The mineralogical analysis further confirms that FA2 contains more minerals with a lower melting point and a low content of mullite.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Quartz</th>
<th>Mullite</th>
<th>Hematite</th>
<th>Magnetite</th>
<th>Amorphous phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>FA1</td>
<td>7.0</td>
<td>39.3</td>
<td>1.2</td>
<td>0.1</td>
<td>39.5</td>
</tr>
<tr>
<td>FA2</td>
<td>7.2</td>
<td>19.1</td>
<td>2.5</td>
<td>3.1</td>
<td>55.2</td>
</tr>
<tr>
<td>FA3</td>
<td>7.8</td>
<td>32.3</td>
<td>1.5</td>
<td>0.2</td>
<td>58.1</td>
</tr>
</tbody>
</table>

As an addition for the experimental work, microsilica (96 % SiO₂) and Fe₂O₃ (powder, 97 % cleanliness) were selected. As a reference, samples from pure fly ash were used, which were then modified with a 5 % or 10 % addition. The mixtures were mixed with water to reach the limit of fluidity. Samples of size 20 mm × 20 mm × 100 mm were made, which were next day dried at 60 °C for 2 h and then fired in a muffle kiln. The firing was characterized by an initial temperature of 25 °C and the rate of firing the muffle kiln of 300 °C/h and an isothermal dwell at 1150 °C (resp. 1200 °C) for 10 min. After firing and natural cooling, the specimens were taken out of the kiln and placed in a desiccator. After thermal stabilization, their density (CSN EN 1015-10), compressive strength (CSN EN 14617-15) and water-absorbing capacity (CSN EN 1097-6) were determined. Then, the samples were analyzed with a scanning electron microscope using a sensing element in an environmental form. Primarily, the structure was analyzed and the influence of the addition on the fly ash body.

3 RESULTS AND DISCUSSION

After firing in a laboratory furnace and sufficient cooling to laboratory temperature 20±5 °C, the following
physico-mechanical parameters of the test specimens were determined.

Figure 1 shows the results of the determination of density. In general, it can be stated that the density grows slightly with an increasing temperature of firing. The samples based on FA1 showed the highest values of density, while the samples based on FA3 showed the lowest density.

A determination of the compressive strength (Figure 2) showed that the strengths at the firing temperature 1200 °C were considerably higher than the strengths at 1150 °C. The addition of silica had a very positive effect on the strength of the fly ash body – samples with as little as 5% showed considerably higher strengths. In contrast, the addition of Fe₂O₃ weakened the fly-ash body and the measured strengths were often lower than those of the reference samples based on pure fly ash.

An evaluation of the results of the determination of the water-absorbing capacity (Figure 3) in some cases shows the influence of firing temperature on the quality of the fly-ash body. The water-absorbing capacity of the samples fired at 1150 °C was often much higher. This can be caused by insufficient sintering of the fly-ash body and a higher proportion of open porosity. The influence of the type of addition on the results of the water-absorbing capacity is most marked for the samples based on FA with the addition of Fe₂O₃.

To clarify the results determined during the evaluation of the physico-mechanical parameters, the samples

Figure 1: Density of fired samples
Slika 1: Gostota žganih vzorcev

Figure 2: Compressive strength of fired samples
Slika 2: Tlačna trdnost žganih vzorcev

Figure 3: Water absorption of fired samples
Slika 3: Absorpcija vode žganih vzorcev

Figure 4: Structure of fly-ash body with 5% of mass fractions of Fe₂O₃
Slika 4: Struktura kosa letečega pepela s 5% masnega deleža Fe₂O₃

Figure 5: Detail of structure of fly-ash body with 5% of mass fractions of Fe₂O₃
Slika 5: Detalj strukture kosa letečega pepela s 5% masnega deleža Fe₂O₃
were examined under a scanning electron microscope with a sensing element in an environmental form. The aim was an evaluation of the influence of firing temperature and additions on the structure of the fly-ash body.

Figures 4 to 7 show the structure of a surface of a test specimen based on FA3 with a 5% addition of Fe₂O₃ and fired at 1200 °C. Figure 4 shows a low proportion of sintered structure and visible grains of fly ash. Figure 5 shows a part of a sample with melted material at higher magnification, where non-reacted grains of Fe₂O₃ are visible. Figure 6 shows a closer detail of the same place grains of Fe₂O₃ and Figure 7 an analysis of the elements in a larger area where the Fe is evident in the grains. This fact proves that Fe₂O₃ did not take part in the formation of a solid structure, and that it weakened the fly-ash body.

Figures 8 and 9 show the structure of the test specimens based on FA3 with a 10% addition of silica. Figure 8 shows a sample fired at 1150 °C, where the grains of fly ash and a minimal proportion of melted material are clear. Figure 9 shows a sample fired at 1200 °C, where the proportion of melted material is more con-
siderable and the grains have a stronger and more homogeneous structure.

4 CONCLUSIONS

The manufacture of artificial aggregate from fly ash is one of the options for using the maximum proportion of this raw material in construction materials. The character of the fly ash and good control of the technology for producing the aggregate by self-firing also brings savings for traditional raw materials. The paper described the experimental testing of possibilities for the modification of fly ash with the addition of microsilica or Fe₂O₃ in order to achieve better quality of the fly-ash body with a lower energy demand of the process. The results show how important it is to set the firing temperature at 1200 °C, which makes sure it is possible to achieve considerably higher strengths in the system. An evaluation of the influence of the used additions showed that the addition of microsilica unambiguously improves the quality of the fly-ash body. Higher strengths and a lower water-absorbing capacity were achieved compared to use of pure fly ash. The addition of Fe₂O₃ did not take part in the formation of the melt, and stayed almost unchanged and weakened the fly-ash body structure.

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