Cost-efficient one-part alkali-activated mortars with low global warming potential for floor heating systems applications

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Increasing building energy efficiency is one the most cost-effective ways to reduce emissions. The use of thermal insulation materials mitigates heat loss in buildings, therefore minimising heat energy needs. In recent years, several papers were published on the subject of foam alkali-activated cements with enhanced thermal conductivity. However, on those papers cost analysis was strangely avoided. This paper presents experimental results on one-part alkali-activated cements. It also includes global warming potential assessment and cost analysis. Foam one-part alkali-activated cements cost simulations considering two carbon dioxide social costs scenarios are also included. The results show that one-part alkali-activated cements mixtures based on 26%OPC + 58.3%FA + 8%CS + 7.7%CH and 3.5% hydrogen peroxide constitute a promising cost-efficient (67 euro/m³), thermal insulation solution for floor heating systems. This mixture presents a low global warming potential of 443 KgCO₂eq/m³. The results confirm that in both carbon dioxide social cost scenarios the mixture 26 OPC + 58.3 FA + 8 CS + 7.7 CH with 3.5% hydrogen peroxide foaming agent is still the most cost efficient.

Keywords: foam agents; alkali-activated cements; compressive strength; thermal conductivity; global warming potential; cost efficiency

1. Introduction

The increasing worldwide demand for energy, is a major cause of unsustainable development on our Planet. Between 2007 and 2030, energy demand will have risen around 40%, reaching a total of 16.8 billion tonnes of equivalent petroleum-TEP (Pacheco-Torgal & Jalali, 2011). The rise in energy consumption has two main causes, the increase in world population and the fact that there are an increasing number of people with access to electricity. In fact, currently, there are 1.5 billion people that still have no access to electricity (UN, 2010). Besides, since the urban human population will almost double, increasing from approximately 3.4 billion in 2009 to 6.4 billion in 2050 (WHO, 2014), this will dramatically increase electricity demand. Since buildings consume throughout their life cycle, more than 40% of all energy produced (OECD, 2003), this subsector has a high potential for reducing carbon dioxide emissions. The European Union has been producing legislation in the field of building’s energy performance, which was materialised in the form of the European Energy Performance of Buildings Directive 2002/91/EC (EPBD), which in turn, has been recast in the form of the 2010/31/ EU by the European Parliament on 19 May 2010. One of the new aspects of the
EPBD that reflects an ambitious agenda on the reduction of the energy consumption is the introduction of the concept of nearly zero-energy building (Pacheco-Torgal, Cabeza, Mistretta, Kaklauskas, & Granqvist, 2013). The use of thermal insulation materials constitutes the most effective way of mitigating heat loss in buildings, thus reducing heat energy needs, and therefore contributing to the nearly zero energy target. These materials are very important for the building material industry and represent a 21 billion € market share (Pacheco-Torgal, 2014). However, most current insulation materials are associated with negative impacts in terms of toxicity. Polystyrene, for example contains anti-oxidant additives and ignition retardants, additionally, its production involves the generation of benzene and chlorofluorocarbons. On the other hand, polyurethane is obtained from isocyanates, which are widely known for their tragic association with the Bhopal disaster. Besides, they release toxic fumes when subjected to fire (Pacheco-Torgal, Jalali, & Fucic, 2012). In terms of legislation, the European Union recently approved the Regulation (EU) 305/2011 (2011) related to the Construction Products Regulation that will replace the current Directive 89/106/CEE, already amended by Directive 1993/68/EEC, known as the Construction Products Directive. Investigations in the field of geopolymers had an exponential increase after the research results of Davidovits (1979) who developed and patented binders obtained from the alkali-activation of metakaolin, having named it after the term “geopolymer” in 1978. For the chemical designation of the geopolymer, Davidovits suggested the name “polysialates”, in which Sialate is an abbreviation for aluminosilicate oxide. However, Provis and Van Deventer (2009) mentioned that the sialate nomenclature “implies certain aspects of the geopolymer gel structure which do not correspond to reality”. Despite all the investigations published on these materials in the last decades, some aspects still needed to be further investigated especially concerning durability performance (Pacheco-Torgal, Abdollahnejad, Camões, Jamshidi, & Ding, 2012; Pacheco-Torgal, Labrincha, Leonelli, Palomo, & Chindaprasirt, 2014; Provis, 2013). In the last years, several papers were published on foam alkali-activated cements with enhanced thermal conductivity (Feng et al., 2015; Hlaváček, Šmilauer, Škvára, Kopecký, & Šuc, 2015; Sanjayan, Nazari, Chen, & Nguyen, 2015; Zhang, Provis, Reid, & Wang, 2014). However, cost analysis was oddly avoided. Those authors strangely mentioned that low-cost alkali-activated cements materials could be produced. However, it is known that one of the shortcomings of alkali-activated cements is its non-cost efficiency. Also Abdollahnejad, Pacheco-Torgal, and Félix (2015) recently studied foam fly ash-based two-part (NaOH, NaSiO₃) alkali-activated cements having report that the mixtures cost more than 300 euro/m³. This means that investigations concerning the development of low-cost foam alkali-activated cements are still needed. The discovery of one-part alkali-activated cements is considered a key event on the evolution of low-carbon alkali-activated cements technology in the “just add water” concept. Contrary to ordinary Portland cement (OPC)-based materials, alkali-activated cements use caustic activators that makes the handling and application of alkali-activated cements very difficult. That is why the alkali-activated cements based on the OPC concept of just add water to a powder are an important research line. However, investigations made so far on one-part alkali-activated cements show that these materials are associated with very low compressive strength (Koloušek et al., 2007; Peng, Wan, Shen, & Xiao, 2014). Some authors even report a compressive strength decrease with time for one-part alkali-activated cements based on calcined red mud and sodium hydroxide blends (Ke, Bernal, Ye, Provis, & Yang, 2015). Abdollahnejad, Hlavacek, Miraldo, Pacheco-Torgal, and Aguiar (2014) recently investigated one-part alkali-activated cements having obtained relevant compressive strength using fly
ash and minor amounts of OPC. Since supply chain risks can limit the alkali-activated cements technology's wider adoption, (Van Deventer, Provis, & Duxson, 2012) the use of minor volumes of OPC can help overcome this problem. Yang and Lee (2013) and Yang, Lee, Song, and Gong (2014) studied foam-ground granulated blast-furnace slag concrete produced with three alkali activators: Ca(OH)₂ plus Mg(NO₃)₂, Ca(OH)₂ plus 6.5% Na₂SiO₃, and 2.5% Ca(OH)₂ plus Na₂SiO₃. These authors concluded that a unit binder content of approximately 400 kg/m³ is required to achieve the minimum quality requirements specified by the Korean Industrial Standard for floor heating systems (Figure 1). This paper discloses results of a study on one-part foam alkali-activated cements based on fly ash and OPC blends floor heating system applications. It also includes global warming potential assessment and cost analysis.

2. Experimental work

2.1. Materials and mix design

The mixtures under investigation were based on the ones described in the international patent authored by Zheng, Van Deventer, and Duxson (2007) and whose mechanical properties and durability were already previously studied (Abdollahnejad, Pacheco-Torgal, Aguiar, & Jesus, 2015; Abdollahnejad et al., 2014). The composition of the dry mix in this study was: kaolin, fly ash, OPC, sodium hydroxide (NaOH), calcium hydroxide (Ca(OH)₂), water, and superplasticizer. Mortars were made with a sand having a maximum dimension of 4 mm and a density of 2660 kg/m³. The OPC is of class I 42.5 R type, containing between 95 and 100% of clinker content, a specific weight of 3.15 g/cm³ and a Blaine fineness of 3842 cm²/g (Table 1). The superplasticizer (SP) used was SIKA 3002 HE. The referred SP was used to maintain a uniform consistency throughout the different mixes. The chemical composition of the fly ash complies with the minimum requirements indicated in EN-450-1 (2012) for use as a partial replacement of cement in concrete. Based on this standard, the fly ash was categorized as class B and group N for the loss of ignition and fineness, respectively. Also, the material in question has a specific weight of 2.42 g/cm³ and a chemical composition as shown in Table 1. The kaolin has a BET surface area of 210.000 cm²/g and its particle size is shown in Figure 1. This is crucial information as this parameter influences the dehydroxylation temperature of kaolin. The DTA/TGA curves for kaolin are presented in Figure 2. A well-defined endothermic DTA peak and sharp weight loss in the TGA

Figure 1. Details of sections of typical floor heating system (Peng et al., 2014).
Curve appears between 550 and 600 °C (Figure 3). This loss results from the transition to the amorphous and more reactive metakaolin phase. A mixture of kaolin and sodium hydroxide calcined in a furnace at 650 °C during 140 min is referred as calcined stuff (CS). A mixture of kaolin and sodium hydroxide was calcined in a furnace at 650 °C during 140 min. The cooled mixture was then ground into powder. Three foaming agents were used, namely aluminium powder (0.5–1.5%), NaBO₃ (3–5%) and H₂O₂ (3–5%). Table 2 shows the mix composition of one-part alkali-activated cements mortars.

3. Experimental procedures

Compressive strength was performed on 50 × 50 × 50 mm³ mortar specimens and was obtained for each mixture from an average of 3 cubic specimens. The specimens were tested after 28 days of curing. The specimens were immersed in water at room temperature for 24 h. First, the weight of the specimens was recorded while suspended by a thin wire and completely submerged in water is recorded as Wim (immersed weight). After that, the specimens were removed from water, and placed for 1 min on a wire mesh.
allowing water to drain, then visible surface water was removed with a damp cloth and weight is recorded as Wsat (saturated weight). All specimens were then placed in a ventilated oven at 105 °C for not less than 24 h while allowing for two successive weighings at intervals of 2 h to show an increment of loss not greater than 0.1% of last previously determined weight of the specimen. The weight of the dried specimens was recorded as Wdry (oven-dry weight). The bulk density was assessed using the water saturation method. Thermal conductivity was assessed using an Alambeta instrument developed at the Technical University of Liberec, Czech Republic. During the measurements, the initial temperature of the samples and the base of the Alambeta were kept at 22–24 °C, and the relative humidity was recorded as being in the range of 55–65%. To investigate the effect of the temperature drop on the thermal properties of the samples, the temperature of the measuring head, at first, was 10 °C higher than the environmental temperature. During the second set of measurements, the measuring head temperature was 40 °C higher than the base plate. To assess the crystalline structure of these mixtures, X-ray diffraction (XRD) analysis was employed. Powder XRD was implemented on a Bruker D8 Discover with Cu-Kα radiation (λ = 1.54060 Å) at 40 kV and 40 mA. Each sample was scanned from 5° to 70° at a speed of .04°s–1. The analysis for phase identification was performed using analytical software EVA. For examination through scanning electron microscopy (SEM), the samples were covered with 40-nm film of Au-Pd (80–20 weight %); in a high-resolution sputter coater, 208HR Cressington Company, coupled to a MTM-20 Cressington High Resolution Thickness Controller. Ultra-high resolution field emission gun SEM, NOVA 200 Nano SEM and FEI Company were also used. Backscattering Electron images were achieved through an acceleration voltage of 15 kV. Chemical analyses of samples were performed with the Energy Dispersive Spectroscopy (EDS) technique, using an EDAX Si(Li) detector with an acceleration voltage of 20 kV.

4. Results and discussion

4.1. XRD, SEM and EDS

The results of XRD are presented in Figure 4. A small mullite crystalline phase is detected in all the calcined kaolin-based mixtures. A clear calcite peak is common to the calcined kaolin mixtures. The clear crystalline structure in these mixtures contrasts with the amorphous structures of traditional two-part alkali-activated cements. The different foaming agents (aluminum powder and hydrogen peroxide) seem not to have altered the crystalline phases of the hardened material, the exception being the clear...
calcite silicate oxide in the mixture containing 5% oxygen peroxide. Results of SEM are presented in Figure 5. Also the mixtures have the following compressive strength after 28 days of curing (2.3, 2.2, 7 and 8.7 MPa). The mixtures associated to lower compressive strength show less dense microstructure. This is clearer in the mixture 30 OPC_58.3 FA_4 CS_7.7 CH (5% H2O2) with 5% hydrogen peroxide foaming agent. Concerning the EDS molar ratios (Table 3) several areas (Z2 e Z3 in mixture with 3.5% H2O2 and Z2 in mixture with 0.5% aluminum powder) in the one-part geopolymer mixtures show lower C/S ratio typical of sodium incorporation in the CSH phase into Na-C-S-H gel (Macphee, 1989), this phase usually evolves into compositions with higher calcium and lower aluminum content (with the C–A–S–H gel proposed to be more stable than N–A–S–H, at high pH) (Garcia-Lodeiro, Fernández-Jiménez, & Palomo, 2013).

Table 3. EDS atomic ratios.

<table>
<thead>
<tr>
<th></th>
<th>100 OPC 1.5% AL</th>
<th>30 OPC_58.3 FA_4 CS_7.7 CH</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Z1</td>
<td>Z2</td>
</tr>
<tr>
<td>SiO2/Al2O3</td>
<td>1.41</td>
<td>3.96</td>
</tr>
<tr>
<td>Al2O3/Na2O</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>CaO/SiO2</td>
<td>6.57</td>
<td>3.32</td>
</tr>
<tr>
<td>Na2O/CaO</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MgO/Al2O3</td>
<td>.57</td>
<td>3.84</td>
</tr>
</tbody>
</table>

Figure 3. DTA/TGA curves for kaolin.
4.2. Compressive strength

Compressive strength results for one-part alkali-activated mortars are shown in Figure 6. For mortars based only in Portland cement, the increase of the aluminium content leads to a decrease in compressive strength except when the aluminium content increases from 1.2 to 1.5%, which leads to the increase of compressive strength. The reaction of aluminium powder in the alkaline environment liberates $\text{Al}_2\text{O}_3$ and hydrogen gas ($\text{H}_2$) according to Equation (1) (Arellano Aguilar, Burciaga Díaz, & Escalante García, 2010):

$$8\text{Al} + 2\text{OH}^- + 2\text{H}_2\text{O} \rightarrow 4\text{Al}_2\text{O}_3^- + 3\text{H}_2$$

When the NaBO$_3$ is used as foaming agent in Portland cement mortars, the compressive strength decreases with the increase in the foaming agent content. Although one exception is detected, when NaBO$_3$ increased from 3 to 3.5% the compressive strength increases. Concerning the use of H$_2$O$_2$, the increase of foaming agent shows a minor increase in the compressive strength but when the foaming agent is increased from 4% to 4.5%, the compressive strength decreased. Comparing the performance of the different foaming agents on Portland cement mortars it is observed that the higher compressive strength (20.24 MPa) was recorded for a .5% AL content. Concerning the mixture 30 OPC_70 FA, the increase of the aluminium content results in a decrease in compressive strength except when its increase from 1 to 1.2% led to slight compressive strength increase. When the NaBO$_3$ is used, a trend on compressive strength decrease with foaming agent increase can be found. With respect to the use of H$_2$O$_2$, the increase in

![Figure 4. XRD patterns of: (a) 100 OPC (1.5% AL); (b) 30 OPC_58.3 FA_4 CS_7.7 CH (.5% AL); (c) 30 OPC_58.3 FA_4 CS_7.7 CH (5% H$_2$O$_2$); (d) 30 OPC_58.3 FA_4 CS_7.7 CH (3.5% H$_2$O$_2$); M – Mullite; Q – Quartz; C – Calcite; CA – Calcite silicate oxide.](image-url)
Figure 5. SEM images of: (a) 30 OPC_58.3 FA_4 CS_7.7 CH (3.5% H₂O₂); (b) 30 OPC_58.3 FA_4 CS_7.7 CH (5% H₂O₂); (c) 30 OPC_58.3 FA_4 CS_7.7 CH (.5% AL); (d) 100 OPC (1.5% AL).
foaming agent from 3 to 3.5% leads to a decrease of the compressive strength. In an alkaline environment, oxygen peroxide decomposes into water and oxygen according to Equations (2) and (3) (Damjanovic, Genshaw, & Bockris, 1967):

\[
H_2O_2 + OH^- \rightarrow 4Al_2O^- + H_2O \quad (2)
\]

\[
HO_2^- + H_2O_2 \rightarrow H_2O + O_2 + OH^- \quad (3)
\]

In general, it can be stated that, the results gathered for the compressive strength of the mixture 30 OPC_70 FA with different foaming agents are taking place in a lower range (in comparison to other compressive strengths of one-part alkali-activated mortars). Regarding the mixture 30 OPC_58.3 FA_4 CS_7.7 CH an increase of the aluminium content results in a reduction in compressive strength. Increasing NaBO_3 from 1 to 5%

<table>
<thead>
<tr>
<th>Constituents of mortars</th>
<th>GWP (kg CO_2eq)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ordinary Portland cement (OPC)</td>
<td>$8.44 \times 10^{-1}$</td>
</tr>
<tr>
<td>Superplasticizer</td>
<td>$7.49 \times 10^{-1}$</td>
</tr>
<tr>
<td>Fine sand, aggregates</td>
<td>$2.40 \times 10^{-3}$</td>
</tr>
<tr>
<td>Free water</td>
<td>$1.55 \times 10^{-4}$</td>
</tr>
<tr>
<td>Metakaolin</td>
<td>$9.24 \times 10^{-2}$</td>
</tr>
<tr>
<td>Fly ash</td>
<td>$5.26 \times 10^{-3}$</td>
</tr>
<tr>
<td>Soda powder</td>
<td>$2.24 \times 10^{0}$</td>
</tr>
<tr>
<td>Sodium silicate</td>
<td>$1.76 \times 10^{0}$</td>
</tr>
<tr>
<td>Ca(OH)_2 – hydraulic lime</td>
<td>$4.16 \times 10^{-1}$</td>
</tr>
<tr>
<td>Oxygen peroxide</td>
<td>$8.01 \times 10^{0}$</td>
</tr>
<tr>
<td>Sodium perborate</td>
<td>$3.77 \times 10^{0}$</td>
</tr>
<tr>
<td>Aluminum powder</td>
<td>$1.23 \times 10^{0}$</td>
</tr>
</tbody>
</table>
reduced the compressive strength with some exceptions. The exceptions were found when the NaBO$_3$ content increased from 1 to 2% and 4.5 to 5%. No specific trend was observed in the compressive strength of the mixture 30 OPC_58.3 FA_4 CS_7.7 CH that used H$_2$O$_2$. Comparing the performance of the different foaming agents, the higher compressive strength (7 MPa) took place for a .5% AL content. Additionally, two similar compressive strengths recorded for this mixture: one with 1% AL (6 MPa) and other with 1.2% AL (6.04 MPa). Results of the compressive strength in mixture 26 OPC_58.3 FA_8 CS_7.7 CH show that there is an increase in compressive strength when the Al content increased. The increase in compressive strength can be due to an increase in density related to pore collapses, as suggested by authors (Masi, Rickard, Vickers, Bignozzi, & van Riessen, 2014). Those authors also confirm that aluminium powder is very reactive, thus requiring a much smaller amount (than hydrogen peroxide) for obtaining similar densities. However, since the former costs 33 times more than the latter (Table 4), it is important to compare the economic performance of the different mixtures (Section 6). Nevertheless, an exception was observed when the aluminium content increased from .5 to 1%, since, no specific trend was observed in compressive strength of the mixture 30 OPC_58.3 FA_4 CS_7.7 CH that used NaBO$_3$. The higher compressive strength (6.99 MPa) was registered for a 4% NaBO$_3$ content. Additionally, concerning the mixture 18 OPC_58.3 FA_16 CS_7.7 CH, an increase in the aluminium content leads to a decrease in compressive strength. When the NaBO$_3$ is used as a foaming agent, a trend on compressive strength increase with foaming agent increase can be noticed, except when the NaBO$_3$ increase from 4 to 4.5% leading to a reduction in compressive strength. With respect to the use of H$_2$O$_2$, the increase of the foaming agent results in a reduction of the compressive strength. By comparing the performance of the different foaming agents, it can be concluded that the highest compressive strength (6.11 MPa) was measured for a 5% NaBO$_3$ content. The overall results show that several mixtures show a compressive strength similar to the one presented by commercial cellular concrete blocks (6.5 MPa). However, most mixtures comply with the compressive strength requirement ($\geq 1.4$ MPa) of the grades 4 and 5 of the Korean Industrial Standard code for foam concrete for floor heating systems (Yang et al., 2014).

4.3. Water absorption by immersion

The results of water absorption of one-part alkali-activated mortars are shown in Figure 7. For mortars based only in Portland cement, the increase of the aluminium content does not lead to a relevant change in water absorption. In fact, when it increases from 1.2 to 1.5%, the water absorption decreases slightly. When NaBO$_3$ is used as a foaming agent in Portland cement mortars, a trend on water absorption increase with foaming agent content increase can be noticed. However, some exceptions are detected such as the rise from 3% to 3.5% and the increase from 4 to 4.5% that lead to a reduction in water absorption. Concerning the use of H$_2$O$_2$, the increase of foaming agent from 3 to 3.5% shows a minor growth in the water absorption, but when the foaming agent is increased above 3.5% no water absorption increase is detected. Comparing the performance of the different foaming agents on Portland cement mortars, it can be seen that the higher water absorption (62%) took place for a 5% NaBO$_3$ content. However, this mixture shows a foaming efficiency (water absorption/foaming agent content = 12) that is lower than that of other mixtures based on an aluminium foaming agent. These mixtures show higher water absorption levels with just .5% content (efficiency = 90). Concerning the mixtures 30 OPC_70 FA, the increase of the aluminium content leads to
a slight increase in water absorption. When the aluminium increases from 1.2% to 1.5% a slight water absorption decrease was detected. When the NaBO₃ is used as foaming agent in this mixture, a trend on water absorption decrease with foaming agent increase was registered. However, there was an exception. When the NaBO₃ increased from 4 to 4.5%, this led to a water absorption reduction. Regarding the use of H₂O₂, the increase of foaming agent depicts a minor decrease in the water absorption. Yet, when the foaming agent is increased above 4%, a slight increase in water absorption is detected. When comparing the performance of the different foaming agents on the mixture 30 OPC_70 FA, it is revealed that the higher water absorption (50.53%) was recorded for 1.5% AL content. With respect to the mixtures 30 OPC_58.3 FA_4 CS_7.7 CH, an increase of the aluminium content leads to an increase of water absorption. On the other hand, the increase from .8 to 1% and 1.2 to 1.5% decreases the water absorption. When the NaBO₃ is used as foaming agent, a decrease was detected by increasing the foaming agent content, although some exceptions were detected from 1 to 2% and 4 to 5% (which led to water absorption increase). Concerning the use of H₂O₂, the increase of the foaming agent leads to a reduction in the water absorption. One exception was observed concerning the increase of the foaming agent from 4 to 4.5% which lead to an increase in water absorption. Comparing the performance of the different foaming agents it is found that the higher water absorption (81.07%) took place for a 4.5% H₂O₂ content. This corresponds to an efficiency of (81.07/4.5 = 18) while Al mixtures show high efficiencies (40/1.2 = 33.3). Concerning the mixtures 26 OPC_58.3 FA_8 CS_7.7 CH, an increase of the aluminium content leads to a slight decrease in water absorption. An exception was noticed when the use of AL increased from .5 to 1% and a minor increase in the water absorption was also detected. When the NaBO₃ is used as the foaming agent the water absorption is almost constant, regardless of the increase on the foaming agent content. With respect to the use of H₂O₂, the increase of the foaming agent shows no change in the water absorption. The performance of different foaming agents on this mixture showed that the higher water absorption (72.72%) was registered for a 3% NaBO₃ content. Additionally, some similarly high water absorptions were observed for this mixture with 1.2% AL (72.31%), 1.5% AL (72.22%), 3.5% NaBO₃.
(72.27%), and 5% NaBO$_3$ (72.70%). Concerning the mixture 30 OPC$_{58.3}$ FA$_{16}$ CS$_{7.7}$ CH, an increase of the aluminium content leads to an increase in water absorption. When the NaBO$_3$ is used, a trend on water absorption reduction with foaming agent content increase was noticed. However, the increase of NaBO$_3$ from 3 to 3.5% and the increase of NaBO$_3$ from 4.5 to 5% lead to a water absorption increase. Regarding the use of H$_2$O$_2$, by increasing foam agent content no increase in water absorption was discovered. Comparing the performance of the different foaming agents on the mixture 30 OPC$_{58.3}$ FA$_{16}$ CS$_{7.7}$ CA it can be seen that the higher water absorption (81.48%) took place for a 5% NaBO$_3$ content.

4.4. Bulk density

The results of the bulk density of one-part alkali-activated mortars are shown in Figure 8. For mortars based only in Portland cement, an increment in the aluminium content leads to a decrease in bulk density. When the NaBO$_3$ is employed in Portland cement mortars, a trend on bulk density decrease with foaming agent content increase can be noticed. Concerning the use of H$_2$O$_2$, the increase of the foaming agent shows a minor increase in the bulk density, but when the foaming agent is increased from 4 to 4.5% slight reduction in bulk density was detected. Regarding the performance of the different foaming agents on Portland cement mortars, it is revealed that the lowest bulk density (1067 kg/m$^3$) was recorded for a 3% H$_2$O$_2$ content. Concerning the mixture 30 OPC$_{70}$ FA the increase of the aluminium content leads to decrease in bulk density except when it increases from 1 to 1.2% leading to a slight bulk density increase. When the NaBO$_3$ is used, a trend can be noticed on bulk density increase. Regarding the use of H$_2$O$_2$, the increase of the foaming agent from 3 to 4% shows a decrease in the bulk density. Yet, when the foaming agent is increased above 4% an increase in the bulk density is registered. The lowest bulk density (785 kg/m$^3$) took place for a 4.5% H$_2$O$_2$ content. In respect to the results of bulk density for mixture 30 OPC$_{58.3}$ FA$_4$ CS$_{7.7}$ CH, increasing the aluminium content leads to a reduction on bulk density, except when

![Figure 8. Bulk density.](image-url)
it increases from .8% to 1% which lead to a slight bulk density increase. Increasing the NaBO$_3$ content results in increasing the bulk density, except when the former increases from 4 to 4.5%, situation that leads to a slight bulk density decrease. Concerning the use of H$_2$O$_2$, the increase of foaming agent content shows a constant trend in bulk density results. Comparing the obtained results by using different foaming agents it is observed that the lowest bulk density (821 kg/m$^3$) took place for a 1.2% AL content. Concerning the mixture 26 OPC_58.3 FA_8 CS_7.7 CH, an increase in the aluminium content from .5 to 1% leads to slight increase in bulk density. On increasing Al content by more than .8%, a constant trend in bulk density (with no increase) was detected. With respect to the increase in NaBO$_3$ content as foaming agent, no change was observed. When NaBO$_3$ content increased from 4 to 4.5% a slight increase in bulk density was noticed. The use of H$_2$O$_2$ showed a slight decrease from 3 to 3.5%, while increasing H$_2$O$_2$ content by more than 3.5% did not show any change in bulk density. Comparing the results obtained from using different foaming agents, it is revealed that the lowest bulk density (980 kg/m$^3$) was measured for a 4% H$_2$O$_2$ content. Concerning the mixture 18 OPC_58.3 FA_16 CS_7.7 CH, the increase of the aluminium content leads to reduction of bulk density. On the other hand, increasing the NaBO$_3$ content, results in a reduction of bulk density. However some exceptions were noticed. Increasing the NaBO$_3$ from 3 to 3.5% and from 4 to 4.5% leads to slight increase in bulk density. The increase in use of H$_2$O$_2$ leads to decrease of bulk density. The lowest bulk density (654 kg/m$^3$) was recorded for a 4.5% H$_2$O$_2$ content.

4.5. Thermal conductivity

Figure 9 presents the results of the thermal conductivity performance of some one-part foam alkali-activated mixtures. Thermal conductivity test were not carried out on all one-part alkali-activated mixtures. Instead, two criteria were considered to select the mixtures for thermal conductivity testing, (a) Mixtures with compressive strength greater than 2 MPa; (b) Mixtures with bulk density lower than 1100 kg/m$^3$. The lowest thermal conductivity performance (.132 W/m K) was recorded for one-part alkali-activated mortars, in mix 30 OPC + 58.3 FA + 4 CS_7.7 CH and 1.2% Al. Furthermore, a close value of thermal conductivity was measured for the same mixture with 1.5% aluminum powder. As the thermal conductivity of commercial autoclaved aerated concrete masonry blocks (Ytong) is around (.17 W/m K), similar thermal conductivity performances can be considered acceptable for thermal insulators like cementitious materials. The mixture 26 OPC + 58.3 FA + 8 CS_7.7 CH with 3.5% hydrogen peroxide shows an acceptable thermal conductivity of .16 W/m.K because it fits the threshold of the grades 4 and 5 of the Korean Industrial Standard code for foam concrete for floor heating systems (Yang et al., 2014).

5. Global warming potential

The global warming potential GWP (KgCO$_2$e) was assessed using the EcoInvent database. Table 4 shows the global warming potential for each mortar constituent. Figure 10 presents the global warming potential-to-thermal resistance ratio. The results show that mixtures with similar thermal conductivity performance have a considerably different global warming potential-to-thermal resistance ratio. This parameter is influenced by the percentage of Portland cement. Higher percentages are associated with higher ratios of global warming potential-to-thermal resistance ratios. Even the mixtures containing
sodium hydroxide have a lower ratio because although the GWP of NaOH is 2.65 times higher than the GWP of Portland cement, the amount of OPC used in those mixtures is much higher. The mixture 26 OPC + 58.3 FA + 8 CS + 7.7 CH with 3.5% hydrogen peroxide foaming has a global warming potential of 443 KgCO$_{2eq}$/m$^3$. 

Figure 9. Thermal conductivity.

Figure 10. Global warming potential to thermal resistance ratio.
The cost of materials are listed in Table 5, which shows that the Al powder foaming agent has a much higher cost than the other two foaming agents. The cost of the mixtures is presented in Figure 11. The results show that using Al powder leads to a significant increase of the cost of the alkali-activated mixtures. The most cost-efficient mixtures (67 euro/m³) were obtained for 26 OPC + 58.3 FA + 8 CS + 7.7 CH with 3.5% hydrogen peroxide foaming agent. The use of mixtures based on the aluminium powder is not feasible because they are not cost efficient at all. The mixture based on the aluminium powder foaming agent with better economic performance (30 OPC + 58.5 FA + 4 CS + 7.7 CH) costs more than 200 euro/m³. This represents more than twice the cost of Ytong masonry blocks (70–80 euro/m³). It is worth remembering that foam mortars based on classical two-part alkali-activated mortars present a cost above 300 euro/m³ (Abdollahnejad et al., 2015). Figure 11 also includes cost simulations for two scenarios: (a) Carbon social cost of 34.7 euro/ton as per US study (Stanford Report, 2015); (b) Carbon social cost of 206.3 euro/ton (Moore & Diaz, 2015). When the cost of the mixtures includes the cost of carbon dioxide emissions mixtures based on hydrogen peroxide, these mixtures show a substantial increase in their cost. A minor increase is noticed in aluminium powder-based mixtures. The results confirm that in both scenarios the mixture 26 OPC + 58.3 FA + 8 CS + 7.7 CH with 3.5% hydrogen peroxide as the foaming agent presents the lowest cost alternative.

Table 5. Costs of the materials (euro/kg).

<table>
<thead>
<tr>
<th>Sand</th>
<th>Cement</th>
<th>Calcium hydroxide</th>
<th>Fly ash</th>
<th>Kaolin</th>
<th>Sodium hydroxide</th>
<th>Water</th>
<th>SP</th>
<th>H₂O₂</th>
<th>NaBO₃</th>
<th>AL</th>
</tr>
</thead>
<tbody>
<tr>
<td>.02</td>
<td>.1</td>
<td>.3</td>
<td>.03</td>
<td>.29</td>
<td>.85</td>
<td>.1</td>
<td>.82</td>
<td>.98</td>
<td>1.5</td>
<td>32.4</td>
</tr>
</tbody>
</table>

Figure 11. Cost of one-part geopolymer mixtures (Gray bars) Materials cost (Red bars) Materials cost for a carbon cost in scenario 1 (Green bars) Materials cost for a carbon cost in scenario 2.
peroxide foaming agent is still the most cost efficient. The results of the cost-to-thermal resistance ratio are presented in Figure 12. The best ratio was detected for the mixture 26 OPC + 58.3 FA + 8 CS + 7.7 CH and is around 10 euro/(m K/W).

7. Conclusions

Buildings are responsible for more than 40% of the energy consumption and greenhouse gas emissions. Thus, increasing building energy efficiency is one the most cost-effective ways to reduce emissions. The use of thermal insulation materials could constitute the most effective way of mitigating heat loss in buildings by minimising heat energy needs. The discovery of one-part alkali-activated cements is considered a key event in the evolution of low carbon alkali-activated technology. Although, they are associated with very low compressive strength, they also show the potential to be used in thermal insulation applications. The results show that the use of aluminium powder is very effective in obtaining foam materials with low thermal conductivity.

An increase of bulk density with the rise of aluminium powder content was noticed and can be related to pore collapses. Aluminium powder allows the production of mortar mixtures with a compressive strength of above 6 MPa, however, its high cost means they are commercially useless when facing the competition of commercial cellular concrete. The mortar mixture based on aluminium powder mortar with the lowest cost (30 OPC + 58.3 FA + 4 CS + 7.7 CH) costs more than 200 euro/m³. This represents more than twice the cost of Ytong masonry blocks (70–80 euro/m³). It is worth remembering that foam mortars based on classical two-part alkali-activated mortars have a cost above 300 euro/m³, which shows that two-part alkali-activated cements are not a cost-effective solution for the production of thermal insulation materials. The mixture 26 OPC + 58.3 FA + 8 CS + 7.7 CH with 3.5% hydrogen peroxide shows an acceptable compressive

Figure 12. Cost-to-thermal resistance ratio.
strength and thermal conductivity complying with the requirements for the grades 4 and 5 of the Korean Industrial Standard code for foam concrete for floor heating systems (≥1.4 MPa and ≤0.16 W/m K). This mixture is cost efficient (67 euro/m³) and has a low global warming potential of 443 KgCO₂eq/m³. When the cost of the mixtures includes the cost of carbon dioxide emissions, mixtures based on hydrogen peroxide show a substantial increase in their cost. A minor increase is noticed in aluminium powder-based mixtures. The results confirm that in both carbon dioxide social cost scenarios the mixture 26 OPC + 58.3 FA + 8 CS + 7.7 CH with 3.5% hydrogen peroxide foaming agent is still the most cost efficient.

Disclosure statement
No potential conflict of interest was reported by the authors.

References


