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Can Portland cement be replaced by low-carbon alternative materials? A study on the thermal properties and carbon emissions of innovative cements

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ABSTRACT

One approach to decarbonising the cement and construction industry is to replace Portland cement systems with lower carbon alternatives that have suitable properties. We show that seven cementitious binders comprised of metakaolin, silica fume and nano-silica have improved thermal performance compared with Portland cement and we calculate the full CO$_2$ emissions associated with manufacture and transport of each binder for the first time. Due to their high porosity, the thermal conductivity of these novel cements is 58–90% lower than Portland cement, and we show that a thin layer (20 mm), up to 80% thinner than standard insulating materials, is enough to bring energy emissions in domestic construction into line with the UK 2013 Building Regulations. Carbon emissions in domestic construction can be reduced by c. 20–50% and these cementitious binders are able to be recycled, unlike traditional insulation materials.

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1. Introduction

Portland cement is one of the most manufactured materials in the world. Over 3 billion tonnes of cement were manufactured in 2012 (Imbabi et al., 2012), and global demand is expected to increase due to the rapid infrastructural development of emerging economies (Schneider et al., 2011; Benhelal et al., 2013). Indeed, global cement production is forecast to reach 3.7–4.4 billion tonnes by 2050, as stated by the World Business Council for Sustainable Development (WBCSD) report in 2009 (Schneider et al., 2011; Benhelal et al., 2013). Cement is primarily used by the construction and geotechnical industries, but there are other emerging applications, including nuclear waste containment, biological and dental ceramics, and water filtration. Cement clinker is produced by calcining limestone (or marl or chalk) with some clay in a furnace at c. 1500 °C and is a significant source of greenhouse gas emissions (GHG), which are usually expressed as CO$_2$ equivalent (CO$_2$eq) and sometimes referred to as "embedded carbon" (Salas et al., 2016). Approximately 900 kg of CO$_2$eq is released per tonne of cement produced by current practices (Hasanbeigi et al., 2010). Thus, the cement industry is estimated to have contributed 5–7% of global anthropogenic CO$_2$ emissions in 2009 (Turner and Collins, 2013; Hienola et al., 2017). The direct release of CO$_2$ from calcination during clinker production is responsible for c. 50% of the emissions from cement manufacture (Fig. 1). Much of the remaining emissions come from the combustion of fossil fuels for calcination, plus excavation, transportation, milling and grinding processes. Given the global effort to curb CO$_2$ emissions in an attempt to mitigate dangerous climate change effects (Hienola et al., 2017) (for example, the 2015 Paris Agreement, a framework for an internationally coordinated effort to tackle climate change), and the expected rise in global demand for cement, reducing emissions from cement manufacture presents an important challenge. As such, the ‘decarbonisation’ of cement production is becoming a more prominent issue for the cement sector, as evidenced by the World Business Council for Sustainable Development International Energy Agency (IEA) Cement Roadmap (2009), the Industrial Decarbonisation & Energy Efficiency Roadmaps to 2050 report and the British Cement Association (BCA73 Carbon Strategy 2005) (World Business Council for Sustainable Development, 2009 and 2011, Industrial Decarbonisation and Energy Efficiency Roadmaps, 2015).

Researchers and industry have focused their attention on using alternative fuels in place of conventional fossil fuels (and so reducing the GHG emissions of the traditional manufacturing process), and developing alternative materials by partially
replacing clinker used in cement systems with waste or by-products (Kajaste and Hurme, 2016). Clinker replacements that have been developed or tested include reused waste, such as industrial by-products like fly ash, or biomass wastes like rice husk ash. Novel binders such as geopolymers or alkali-activated cement (Turner and Collins, 2013; Rostami and Brendley, 2003; Cruz-Yusta et al., 2011; Wu et al., 2015; Sturm et al., 2016; Nie et al., 2016) have also gained popularity. In most cases, clinker is only partially replaced (Hemalatha et al., 2016) but it produces a ‘greener’ cement. This is advantageous from a regulatory perspective since the existing standardised codes of practice for Portland cement can be adapted or built upon. It is important that the mechanical properties of alternative cements are similar to (or more advantageous than) the properties of Portland cement. Other additives such as silica fume and nano-silica particles improve the properties of Portland cement (Yu et al., 2000; Sanchez and Ince, 2009; Aggarwal et al., 2015; Lazaro et al., 2016) and metakaolin-based geopolymers (Gao et al., 2013; Villaquiran-Caicedo et al., 2015). The main reason geopolymer binders have not yet been more widely adopted by industry is the current lack of regulatory standards backed by long term testing and development (Heidrich et al., 2015).

Portland cement is used in the preparation of mortar for wall rendering/finishing and also in aerated concrete blocks employed as a thermal insulation material (Ahmed and Fried, 2012; Zhang et al., 2014a). However, aerated cement does not offer thermal conductivity values comparable to other solutions on the market, such as polymer foam, glass fibres and vacuum insulation panels (Al-homoud, 2005). Although these materials have very low thermal conductivity, in the range 0.01–0.002 W/(m K) (Cho et al., 2014; Aldawi and Alam, 2016), which can help reduce energy consumption, their production is polluting (Papadopoulos and Giama, 2007; Proietti et al., 2013) plus they cannot entirely be recycled and have to be disposed of in landfill. Geopolymers and cement-free mixtures have been proposed as alternative insulation materials to Portland cement-based composites and have shown thermal conductivity values, 0.17–0.35 W/(m K), lower than traditional cement mortar or concrete (0.2–0.8 W/(m K) (Loudon, 1979)) although not comparable with insulation materials such as glass fibres or polymers (Villaquiran-Caicedo et al., 2015).

Life cycle analyses on selected geopolymer binders have found that their use in place of cement could reduce GHG emissions from the cement industry by 9—64% (Turner and Collins, 2013; McLellan et al., 2011). However, these life cycle emissions are context and country dependent and often subject to availability of the raw materials (Stafford et al., 2016a, 2016b; Moretti and Caro, 2017). To date, the environmental sustainability of a range of cement free mixtures has not been comparatively explored, nor has there been a comprehensive analysis of properties of alternative cements and their potential to completely replace Portland cement. Here we consider the carbon reduction that could be achieved by using seven alternative cementitious materials in place of Portland cement, by evaluating the CO$_2$ eq. gas emissions of Portland cement and geopolymer production and taking the whole life-cycle into account, including the transport of raw materials and the manufacturing process (Imbabi et al., 2012).

The aim of this work is to develop novel ‘green’ cementitious materials with superior thermal properties to Portland cement and low environmental impact. Silica particles, metakaolin and calcium hydroxide are combined in binary or ternary systems and their physical, thermal and mechanical properties are characterised. Thermal performance is calculated in the context of a typical UK domestic construction and a comparison of GHG emissions for these novel cementitious binders and Portland cement is presented for the first time in the UK-European context. These novel Portland cement free binder represent an environmentally friendly alternative with strong potential for recycling, a simple manufacturing process and are able to ensure thermal comfort within current international standards. Furthermore, GHG emissions are calculated following a simplified life cycle assessment methodology, which provides a useful decision-making tool to industries or practitioners to rapidly calculate the carbon footprint of Portland cement free binders.

2. Materials and methods
2.1. Materials

Portland cement samples were prepared using Portland cement CEM I 42.5-R (CAS number 65997-15-1), commercially available from the Lafarge Cement Group, and deionised water (W). Physico-chemical properties of Portland cement are listed in Table S1 of the Supplementary Material. Portland cement were prepared with a liquid to solid (l/s) ratio of 0.3:1 using a rotary mixer according to BS EN 196-1:2016 and cast into cubic moulds for 24 h.
samples were kept for 28 days at relative humidity of 98 ± 2% and temperature of 21 ± 2 °C in a nitrogen gas environment to minimise carbonation prior to testing. Novel Portland cement free samples were prepared using different starting materials. Reagent grade calcium hydroxide, Ca(OH)$_2$ (CAS Number 1305-62-0) and Ludox T50 nano-SiO$_2$ aqueous suspension (CAS number 7631-86-9) were purchased from Sigma Aldrich. Silica fume (CAS number 69012-64-2), commercially available as SF920D from Elkem Microsilica (Norway), was used. Metakaolin was obtained from calcination of kaolin (China clay type purchased from Imerys UK, CAS number 1332-58-7) at 750 °C over 24 h, as described by Alonso et al. (Alonso and Palomo, 2001). Reagent grade sodium hydroxide, NaOH (CAS number 1310-73-2) of nominal concentration 10 M was purchased from Fisher Scientific. Chemical and physical properties of the starting materials (calcium hydroxide (CH), nano-silica (NS), metakaolin (MK), silica fume (SF)) are reported in Table S2 of the Supplementary Material. Given the pozzolanic reactivity of nano-silica and silica fume, binary mixes using calcium hydroxide and silica (nano-silica or silica fume) were investigated (samples CH, CH10, CHNS). Alkali activated binders were prepared mixing metakaolin with calcium hydroxide in different proportions. Sodium hydroxide 10M was added as an activator in samples MK10, AMK, BMK and also CH10, which did not contain metakaolin. Finally metakaolin was mixed with nano-silica and calcium hydroxide, using a lower concentration solution of NaOH (1 M) as the activator (sample MKNS). Mix proportions and sample identification are listed in Table 1. Fresh paste was cast into cubic moulds and specimens were kept for 28 days at relative humidity of 98 ± 2% and temperature of 21 ± 2 °C in a nitrogen gas environment to minimise carbonation. Sample MK10 was thermally prepared following a methodology developed for geopolymers (Zhang et al., 2014b). After mixing, specimens were cast into a cubic mold and kept in an oven at 60 °C and atmospheric pressure for 24 h, then placed in a sealed environment for 28 days at relative humidity of 98 ± 2% and temperature of 21 ± 2 °C.

### 2.2. Physical, thermal and mechanical properties

After ageing for 28 days samples were removed from the mold and dried at 60 °C to remove pore water and perform mechanical tests and micro-structural analyses. Water removal has an impact on the microstructure, therefore analysis and results presented should be regarded comparatively. Compressive strength testing was performed according to BS EN 196-1:2016, using a uniaxial compressive testing machine at a constant strain rate of 0.4 mm/min until fracture (Sanchez and Ince, 2009; Lin et al., 2010). Three specimens of each composite were tested. The resistance value (Rc) is given in MPa as a mean value of three replicates for each mixing. The heat of hydration was measured using an isothermal calorimeter (I-Cal 4000 HPC, Calmetrix). Fresh paste (c. 60 g) was cast into a cylindrical container and placed into the calibrated calorimeter, at a constant temperature of 21 ± 2 °C. The heat flow was recorded over 80 h. Open porosity (p) was calculated using the equations reported in the Methods section of the Supplementary Material. Samples were oven dried at 60 °C to constant mass followed by evacuation in a vacuum chamber then saturated overnight with water in the same chamber.

For each sample the laser flash method (LFA) was used to estimate the coefficient of thermal conductivity (λ), given in W/(m·K). A Netzsch instrument 427 LFA was used. Samples of each composition were tested in an argon atmosphere and thermal conductivity was calculated at 25°C, 60°C and 105°C according to the BS EN 821-2:1997.

The specimens were powdered and pelletized using an hydraulic press to make pellets of 0.127 mm and 3 mm thickness. The surface was coated with graphite to minimise reflectance of the laser beam. A pyroceramic standard supplied by Netzsch was analysed and used as a reference material to calculate the specific heat capacity and thermal diffusivity. Thermal conductivity was calculated at 25°C, 60°C and 105°C, as a function of the open porosity, using the equations reported in the Methods section of the Supplementary Material.

In order to evaluate the insulation properties of these novel cement composites, the thermal transmittance (U) of a typical wall was calculated, using the equations reported in the Methods section of the Supplementary Material. An external wall (1 m high and 1 m wide) of standard UK domestic construction was considered, as shown in Fig. 6 (left). The wall consists (from outdoor to indoor) of horizontal bricks (225×112×65 mm BS EN 771-1:2011, λ = 0.84 W/(m·K)) with a 5 mm layer of cement mortar (λ = 1.4 W/(m·K), Cho et al., 2014) and externally finished with an 18 mm thick layer of mortar render (λ = 1.4 W/(m·K)). Moving inwards from the outer brick skin is a 20 mm thick air cavity (λ = 0.03 W/(m·K)), 9 mm layer of plywood (λ = 0.14 W/(m·K)), a 40 mm thick rockwool insulation wall (λ = 0.04 W/m·K, Al-homoud, 2005) and a 15 mm thick gypsum plaster board (λ = 0.21 W/(m·K)) (Cho et al., 2014), finished with 2 mm thick layer of waterproof plaster paint (λ = 0.09 W/(m·K) (Cho et al., 2014)). This is a pattern in the construction that repeats itself every 70 cm in the vertical direction. Therefore a 1 m wide and 0.7 m high portion of the wall was considered, as it is representative of the entire wall. One-directional heat transfer and constant thermal conductivity values are assumed.

### 2.3. Powder X-Ray diffraction and Scanning Electron Microscopy

Powder XRD analyses were performed using a Bruker D8 Advance diffractometer with CuKα radiation over the range 5–60° 2θ, step size of 0.02° 2θ and 0.5 s/step. DiffracEva software from Bruker was used for XRD pattern evaluation and phase identification. Microstructural analysis of samples was carried out using Scanning Electron Microscopy (W-SEM, Hitachi S-3700N and FE-
2.4. Greenhouse gas emission assessment

Calculation of the total greenhouse gas emission (GHG), expressed as carbon dioxide equivalent (CO$_{2eq}$) per 1000 kg of cement produced, takes into account the collective contribution of CH$_4$, NOx, SOx, CO$_2$ and synthetic gases emitted during production of the material, including excavation and transport of raw materials and reagents, and manufacturing. The approach to estimate the total GHG is based on the methodology reported in McLellan et al. (2011) and calculated using equation (1):

$$GHG_{Tot} = \sum_{i=1}^{n} m_i (d_i e_i + p_i)$$

(1)

where $GHG_{Tot}$ is the total greenhouse gas emission (kg CO$_{2eq}$) per tonne of material produced, $m_i$ is the fraction of component $i$, $d_i$ is the distance transported by a given mode of transport (km), $e_i$ is the emission factor for the transportation mode (kg CO$_{2eq}$/km tonne) and $p_i$ is the emissions per unit mass of component $i$ produced (kg CO$_{2eq}$/tonne). The following assumptions were made in the analysis:

1. The calculations were based on the manufacture of 1 tonne of Portland cement binder and 1 tonne of cement free materials in the United Kingdom, using, where possible, UK products, or otherwise materials from a typical supply chain.
2. Previously published values for CO$_{2eq}$ emissions from the manufacture of the raw materials were used, and added to the emissions from transport to and within the UK.
3. The emissions due to the addition of water to cement paste are very low (0.271 kg CO$_{2eq}$/tonne (Reffold et al., 2008)) and so are not taken into account.
4. Maximum distances and mode of transport are selected as those which maximise CO$_{2eq}$ emissions, because this work adopts the worst-case scenario for CO$_{2eq}$ emissions.
5. Emission factors associated with road transport ($e_r$) and sea transport ($e_s$) are respectively 0.09 kg CO$_{2eq}$/km tonne and 0.02 kg CO$_{2eq}$/km tonne (McLellan et al., 2011; IPCC, 2006).
6. Emissions per unit mass of Portland cement ($P_{PC}$) are 750 kg CO$_{2eq}$/tonne and is produced in mainland UK.
7. Emissions per unit mass of metakaolin ($P_{MK}$), produced in England and silica fume ($P_{SF}$), produced in Norway, are respectively 236 kg CO$_{2eq}$/tonne and 7 $\times$ 10$^{-6}$ kg CO$_{2eq}$/tonne (McLellan et al., 2011; Duxson et al., 2007).
8. The manufacture of calcium hydroxide is based on the hydration of calcium oxide, produced in Northern Ireland, ($P_{CaO} = 750$ kg CO$_{2eq}$/tonne) taking into account a correction factor of 0.97 due to the addition of water ($P_{CH} = 720$ kg CO$_{2eq}$/tonne) as explained in the IPCC Guidelines for national greenhouse gas emissions (McLellan et al., 2011; Duxson et al., 2007).
9. Sodium hydroxide is produced in Northern Ireland by a chemical process using electrolytic cells. The emissions associated with the production are in the range 1120–1915 kg CO$_{2eq}$/tonne as reported for a nominal concentration of 16 M (Turner and Collins, 2013; Mellado et al., 2014; Chan et al., 2016). In order to take into account lower sodium hydroxide concentrations, we used a correction factor of 0.43 and 0.63 respectively for NaOH 1 M and NaOH 10 M on the lowest emission value ($P_{NaOH} = 1120$ kg CO$_{2eq}$/tonne), following the principle of the IPCC guidelines (IPCC, 2006).

10. The nano-silica suspension is manufactured in Germany and the carbon emissions value can be obtained from the manufacture of sodium silicate solution ($P_{NS} = 386$ kg CO$_{2eq}$/tonne) (Lazarro et al., 2013, 2016; EU, 2007).

A schematic diagram of mode of transport and distances for each raw material is shown in Fig. 2.

3. Results and discussion

3.1. Physical, thermal and mechanical properties

The particle size and the high specific surface area of nanoparticles play an important role in the physical and mechanical properties. The measured bulk density ($\rho$), matrix density ($\rho_{mat}$), open porosity ($\varphi$), compressive strength ($R_c$) and cumulative heat released values are reported in Table 2. All the mixes show values of bulk density in the range 600–1100 kg/m$^3$, much lower than standard Portland cement (1900 kg/m$^3$). Density and porosity values are in good agreement with literature data on lightweight materials such as calcium silicate boards and aerated concretes (Hamilton and Hall, 2005; Ünal et al., 2007; Palmero et al., 2015). Sample CHI10 shows a higher bulk density and lower porosity compared to sample CHI due to the greater l/s ratio and the presence of an alkaline activator. Samples MK10, AMK and BMK show very similar density and porosity values and porosity is highest when nanosilica is used. Mechanical tests performed on all the samples after 28 days of curing show values of compressive strength, in the range of 1.8–7.8 MPa. Although compressive
Villaquiran-Caicedo et al. (2015) shows the heat of hydration of C-S-H, which is the most abundant component of hydrated cement paste and responsible for early strength development and hardening (Taylor, 1998) or calcium (sodium) silicate hydrate (C-(N)-S-H) (Gomez-Zamorano et al., 2016; Gomez-Zamorano et al., 2017). Semi-quantitative analysis of the XRD patterns showed that, despite the high pH, sample CHI10 has 54% C-S-H compared to sample CHI (61%). The added Na⁺ concentration requires Ca^{2+} to produce C-N-S-H in addition to the C-S-H produced. Some minor carbonated phases are detected, (calcite and sodium carbonate), arising from surface carbonation. In the mixes containing metakaolin and calcium hydroxide (sample AMK, BMK and MKNS), stratlingite (St), calcium aluminate hydrate (C-A-H) and monocarboaluminate (M) phases are detected, in agreement with Silva et al. 2014. Stratlingite is the main hydrate phase responsible for strength development in lime-metakaolin based materials. An increase of metakaolin content from 25% to 33% respectively in samples AMK and BMK results in well defined peaks of stratlingite, and consequently higher compressive strength.

3.2. Powder X-ray diffraction and Scanning Electron Microscopy

XRD patterns obtained for the developed materials are presented in Figure S2 in the Supplementary Material, where only the major mineral phases are shown. Samples CHI and CHI10 are mainly crystalline portlandite (P) and poorly crystalline calcium silicate hydrate gel (C-S-H), the most abundant component of hydrated cement paste and responsible for early strength development and hardening (Taylor, 1998) or calcium (sodium) silicate hydrate (C-(N)-S-H) (Gomez-Zamorano et al., 2016; Gomez-Zamorano et al., 2017). Semi-quantitative analysis of the XRD patterns showed that, despite the high pH, sample CHI10 has 54% C-S-H compared to sample CHI (61%). The added Na⁺ concentration requires Ca^{2+} to produce C-N-S-H in addition to the C-S-H produced. Some minor carbonated phases are detected, (calcite and sodium carbonate), arising from surface carbonation. In the mixes containing metakaolin and calcium hydroxide (sample AMK, BMK and MKNS), stratlingite (St), calcium aluminate hydrate (C-A-H) and monocarboaluminate (M) phases are detected, in agreement with Silva et al. 2014. Stratlingite is the main hydrate phase responsible for strength development in lime-metakaolin based materials. An increase of metakaolin content from 25% to 33% respectively in samples AMK and BMK results in well defined peaks of stratlingite, and consequently higher compressive strength.

Faujasite (F) is the main crystalline compound in sample MK10 along with C-S-H gel, calcium aluminate hydrate and minor stratlingite. In sample MK10, mixing metakaolin with 10 M NaOH solution promotes alkaline activation and leads to the formation of sodium aluminate silicate hydrate (N-A-S-H) gel and the secondary formation of faujasite (F) (Zhang et al., 2014b; Reig et al., 2016). In sample MKNS, reducing the concentration of the activator from 10 M to 1 M and the addition of calcium hydroxide at ambient temperature results in the precipitation of poorly-crystalline calcium aluminate hydrate (C-A-H), the main phase detected. Sample CHNS presents broad humps at c. 29° and 32° 2θ, typical of C-S-H gel (Garbev et al., 2008).

As shown in the SEM images, the developed materials present a highly porous matrix in agreement with the density and porosity values measured. In sample CHI the matrix is mainly poorly-crystalline C-S-H whereas the presence of NaOH as alkaline activator in sample CHI10 promotes the formation of C-S-H combined with C-(N)-S-H phases, respectively in Fig. 4a and Fig. 4b. As shown in XRD patterns, alkaline-activation of metakaolin/natural lime mixtures results in formation of calcium aluminate silicate hydrate (stratlingite) and C-S-H (sample BMK, Fig. 4c). Fig. 4d shows a semi-crystalline C-S-H phase forming a complex plate-like structure in sample CHNS.

3.3. Thermal conductivity measurements

Thermal conductivity values at 25°, 60° and 105° C calculated according to equation S(5) are shown in Fig. 5 and compared to Portland cement. Values are in the range 0.05–0.26 W/(m K), 50–90% lower than Portland cement. Samples made mixing metakaolin and sodium hydroxide (MK10, AMK and BMK) show thermal conductivity values in accordance with Palmero et al. (2015) and Villaquiran-Caicedo et al. (2015). The addition of silica nano-

<table>
<thead>
<tr>
<th>Sample</th>
<th>ρ (kg/m³)</th>
<th>ρ_{net} (kg/m³)</th>
<th>ϕ</th>
<th>R&lt;sub&gt;c&lt;/sub&gt; (MPa)</th>
<th>Heat release (J/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>OPC</td>
<td>1940</td>
<td>2460</td>
<td>0.21</td>
<td>51.2</td>
<td>235</td>
</tr>
<tr>
<td>CHI</td>
<td>940</td>
<td>2430</td>
<td>0.61</td>
<td>6.4</td>
<td>44</td>
</tr>
<tr>
<td>CHI10</td>
<td>1120</td>
<td>2160</td>
<td>0.48</td>
<td>7.7</td>
<td>211</td>
</tr>
<tr>
<td>MK10</td>
<td>1020</td>
<td>2190</td>
<td>0.53</td>
<td>5.2</td>
<td>446</td>
</tr>
<tr>
<td>AMK</td>
<td>900</td>
<td>2180</td>
<td>0.59</td>
<td>4.7</td>
<td>75</td>
</tr>
<tr>
<td>BMK</td>
<td>850</td>
<td>2020</td>
<td>0.58</td>
<td>6.5</td>
<td>104</td>
</tr>
<tr>
<td>MKNS</td>
<td>640</td>
<td>2260</td>
<td>0.72</td>
<td>1.7</td>
<td>51</td>
</tr>
<tr>
<td>CHNS</td>
<td>610</td>
<td>2390</td>
<td>0.74</td>
<td>2.2</td>
<td>148</td>
</tr>
</tbody>
</table>
particles has a beneficial effect on the thermal conductivity. Sample MKNS and CHNS in fact show the lowest \( \lambda \) values at 25 °C, 0.055 and 0.088 W/(m K) respectively. These values are typical of insulating materials (Cho et al., 2014; Fricke et al., 2008). This effect is attributed to the smaller nano-silica particle size range and greater surface area, which increases the porosity (\( \phi = 0.7 \)) but decreases the pore-size; the overall consequence is an enhanced phonon scattering effect which reduces heat transfer (Alvarez et al., 2010). Samples made by mixing CH and SF, either with water or alkali-activated show a different thermal behaviour: while sample CHI has a thermal conductivity value (\( \lambda = 0.09 \) W/(m K)) similar to CHNS, sample CHI10 has a higher \( \lambda \), suggesting that the alkali-activator (NaOH, 10 M) contributes to the reduction of porosity but decreases the thermal resistance. As shown in XRD patterns, sample CHI contains C-S-H and portlandite, whereas CHI10 is made of C-S-H, natrite and portlandite, bound together in a denser and less porous matrix (c. 20% less than CHI).

Thermal transmittance (U-value) for a typical wall (Fig. 6) was

![Fig. 4. SEM images of (a) sample CHI, (b) sample CHI10, (c) sample BMK, and (d) sample CHNS.](image_url)

**Fig. 4.** SEM images of (a) sample CHI, (b) sample CHI10, (c) sample BMK, and (d) sample CHNS.

![Fig. 5. Thermal conductivity of samples at 25°, 60° and 105°C and porosity values.](image_url)

**Fig. 5.** Thermal conductivity of samples at 25°, 60° and 105°C and porosity values.
calculated to be 0.32 W/(m K), using equations reported in the Methods section of the Supplementary Material. Building Regulation 2013 in England and Wales for refurbishment of existing buildings (domestic and non-domestic use) requires values less than 0.30 W/(m K). The application of a layer of novel cementitious material can contribute to the reduction of the total transmittance below the limit imposed by building regulations, using materials of relatively simple manufacture. The U-value was then calculated taking into account an additional layer of developed material placed in between the bricks and the air cavity. The thickness was chosen in order to minimise the total transmittance below the limit of the building regulations. Thickness values of all the mixes are summarised in Table 3. The thickness of insulation material layers used in the construction industry is in the range of 30–100 mm (e.g. glass fiber, rock-wool or polymeric foam (Cho et al., 2014)). Here, a 20 mm layer of mix MKNS is required to reduce the total transmittance by 10%, as shown in Fig. 6. Conventional insulation materials such as rock-wool, polystyrene or glass fibres, are usually placed in layers of approximately 40–80 mm (Aldawi and Alam, 2016; Bull et al., 2014).

### 3.4. Life cycle emissions

Previous studies have addressed the need to meet thermal requirements, using thermally resistant polymers or composites, but the carbon footprint associated with their manufacture is often overlooked (Fricke et al., 2008; Alam et al., 2014)). The estimated CO$_2$eq emissions (GHG$_i$) for each of the seven cementitious material are reported in Fig. 7 and compared to Portland cement. These present the ‘worst case scenario’, and so the actual CO$_2$eq emissions would likely be lower than those reported here. The carbon footprint of each component material is shown in Fig. 7a. The calculated values are similar to previously published estimates for geopolymer binders and concrete in different contexts Turner and Collins, 2013; McLellan et al., 2011; Mellado et al., 2014, Chan et al., 2016. The results show that all types of novel cements studied here have lower embedded carbon than Portland cement. For example, sample MKNS has the lowest CO$_2$eq emissions associated with its manufacture, estimated to be half the emissions of Portland cement. Sample AMK, which has the highest embedded carbon

<table>
<thead>
<tr>
<th>Sample</th>
<th>Minimum thickness (mm)</th>
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<tbody>
<tr>
<td>CHI</td>
<td>30</td>
</tr>
<tr>
<td>CHI10</td>
<td>80</td>
</tr>
<tr>
<td>MK10</td>
<td>50</td>
</tr>
<tr>
<td>AMK</td>
<td>60</td>
</tr>
<tr>
<td>BMK</td>
<td>80</td>
</tr>
<tr>
<td>MKNS</td>
<td>20</td>
</tr>
<tr>
<td>CHNS</td>
<td>30</td>
</tr>
</tbody>
</table>

Fig. 6. Typical external composite brick wall of domestic building in United Kingdom. (Left: Wall section and temperature (T) profile. Right: Wall-section including a layer of MKNS and temperature profile).

Fig. 7. Total GHG emission and contribution of each raw material for all the mixes. (a) Bubbles indicate the single component in each mix and the size indicates the GHG emission associated: clinker (750 kg CO$_2$eq/tonne), CH (720 kg CO$_2$eq/tonne), SF (0.01 kg CO$_2$eq/tonne), NS (390 kg CO$_2$eq/tonne), MK (236 kg CO$_2$eq/tonne), NaOH 10 M (700 kg CO$_2$eq/tonne), NaOH 1 M (481 kg CO$_2$eq/tonne).
among the novel cements, still has 20% lower CO$_2$eq than Portland cement. The selected raw materials, their world-wide availability coupled with minimum manufacturing make these novel binders environmentally competitive compared to traditional insulators (e.g. 1 tonne of extruded polystyrene is responsible for 1180 kg CO$_2$eq) Papadopoulos and Giama, 2007. NaOH and CH are the most common ingredients of the alternative cements tested here, and the embedded carbon in these materials is similar to clinker (Fig. 7a). Thus, it is the relative proportion of low carbon materials such as SF, MK and NS which determine the overall carbon footprint for each cement. The major energy expended in the manufacture of NaOH occurs in the electrolysis process followed by cooling, which has a large electricity requirement. However, recent findings have proven that natural highly alkaline materials, such as red mud, could be used, with comparatively high mechanical performance Nie et al., 2016. CH is produced by calcination of calcium carbonate followed by hydration. The CO$_2$ footprint of both materials could be reduced if they were produced using an alternative source of energy for the electricity required (e.g. wind turbine, nuclear energy, photovoltaic energy for the manufacture of NaOH) or using biomass or other green fuel in the pyroprocessing of calcium carbonate.

3.5. Environmental impact

Due to their low thermal conductivity, the novel cements present an environmentally sustainable alternative for purposes such as wall cladding. Improving insulation in homes and buildings is an important aspect of reducing thermal energy loss and thus in turn reducing energy consumption. The innovative binders studied here are also highly recyclable compared to conventional insulating components such as polymeric foams, polystyrene, polyurethane, rock-wool or vacuum insulation panels. They could be re-used in the building industry as intended by the European Waste Framework Directive 2008/98/EC and the EU Framework Programme for Research and Innovation Horizon 2020 which stipulates that up to c. 80% of recycled construction and demolition waste material should be re-used to decrease the content of Portland cement used and consequently reduce the amount of waste to be placed in landfill. Further, they require less manufacturing and processing, and the raw materials and reagents are readily available, which is important to consider for large-scale production.

Thus, although novel cements cannot replace Portland cement in all applications, they offer an environmentally sustainable alternative to traditional materials for several applications, and there is significant potential for these materials to contribute towards the decarbonisation of the cement industry.

4. Conclusions

In this study low-carbon cementitious materials have been developed and characterised. Metakaolin, silica fume, nano-silica and calcium hydroxide were combined at different ratios to produce ‘green’ binders for construction industry. Physical and mechanical properties were investigated. Compressive strength values (in the range 2–7 MPa) are typical of non-structural cements (mortars, rendering cements, etc.); density and porosity measurements show that these materials could be used in construction industry as functional building elements. Pozzolanic activity was measured by isothermal calorimetry and hydrated phases (calcium/ aluminum silicate hydrate, faujasite, stratlingite) were found in XRD diffractograms. SEM images give an insight to the microstructure, with the presence of poorly crystalline phases (i.e. C-S-H) and highly porous matrices, in agreement with the porosity measurements (0.48–0.74). Samples present thermal conductivity (0.05–0.26 W/m K), in the range of conventional insulating materials. While previous studies have focused their attention on solely physical properties of Portland cement-free cements and geopolymers, here we have brought together innovative materials able to satisfy thermal performance requirements within environmental standards. In fact, the addition of a 20 mm layer of sample MKNS to an external wall of existing housing, contributes a 10% decrease in thermal transmittance, as required by the Building Regulation 2013 in England and Wales. The environmental impact of these new cements was assessed, including estimating the greenhouse gas emissions associated with their manufacturing and production. All samples have a carbon footprint up to 23–55% lower than Portland cement. They are also more readily made and recycled. These materials are therefore more environmentally sustainable than Portland cement and could help to reduce CO$_2$eq emissions from the cement industry, and reduce heat demand in housing. The life cycle analysis presented here is simplistic, and more detailed life cycle and cost analyses should be the subject of future research to fully understand the economic impact of these materials in replacing Portland cement. However, the methodology adopted provides the basis for implementing a decision-making tool that can advise on, or scope in, low-carbon options before a more resource intensive life cycle assessment approach is applied. It will be therefore useful to construction companies or private developers intending to develop non-conventional building materials (e.g. geopolymers, alkali-activated cements), not yet regulated by law or international standards.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.jclepro.2012.02.138.

References


