Factors influencing the compressive strength of pulverised fuel ash based geopolymers

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**Abstract**

Several factors affecting the reactivity of pulverised fuel ash (pfa) as a precursor for geopolymer concrete have been investigated. These include physical and chemical properties of various pfa sources, inclusion of ground granulated blast furnace slag (ggbs), chemical activator dosages and curing temperature. Alkali-activated pfa was found to require elevated curing temperatures and high alkali concentrations. A mixture of sodium hydroxide and sodium silicate was used and this was shown to result in high strengths, as high as 70 MPa at 28 days. The presence of silicates in solution was found to be an important parameter affecting strength. Detailed physical and chemical characterisation was carried out on thirteen pfa sources from the UK. The most important factor affecting the reactivity was found to be the particle size of pfa. The loss on ignition (LOI) and the amorphous content are also important parameters that need to be considered for the selection of pfa for use in geopolymer concrete. The partial replacement of pfa with ground granulated blast furnace slag (ggbs) was found to be beneficial in not only avoiding the need for elevated curing temperatures but also in improving compressive strengths. Microstructural characterisation with scanning electron microscope (SEM) coupled with energy dispersive X-ray spectroscopy (EDS) was performed on pfa/ggbs pastes. The reaction product of pfa and ggbs in these binary systems was calcium aluminium silicate hydrate gel (C-A-S-H) with inclusion of Na in the structure.

**Keywords:** geopolymer, alkali activated material, fly ash, slag, structural characterisation, mechanical properties, alkali dosages optimisation

# Introduction

The term “geopolymer” was introduced by Davidovits in the 1970s referring to alkali-activated metakaolin [1]. It has since been used for a range of synthetic low-calcium aluminosilicate polymeric materials, as a sub-range of a more general definition which is alkali-activated binders (AAB). Although the initial studies were focused on geological materials such as metakaolin activated with siliceous solutions [2-6], the potential of using other synthetic reactive aluminosilicate materials activated with a range of concentrated alkaline solutions became apparent [1,7-8].

AABs have been studied for the last 40 years [9] and have applications in ceramics, hazardous waste containment, fire-resistant construction materials and refractories [10, 11]. One of the most interesting applications is their use as a cement-free binder that can replace Portland cement-based pastes in construction materials such as concrete and mortar products [12].

Geopolymers can provide a desirable alternative to Portland cement (PC) binders, not only for the environmental benefits arising from the avoidance of CO2 emissions associated with PC production, but also in terms of their performance and durability, where such properties are not only equivalent, but often better than those achieved with PC. The nature of the reactions occurring in geopolymerisation can be summarized in three basic steps [13]:

1. *Dissolution of the aluminosilicate solids:* aluminosilicates in the pozzolanic solid are dissolved by alkaline hydrolysis in the high pH solution of the concentrated alkaline activator. This forms a solution of silicate, aluminate and aluminosilicate species.
2. *Gel Formation:* species released by dissolution are held in the aqueous phase, which may also contain silicate present from the activating solution. This supersaturated aluminosilicate solution forms a gel as oligomers form long chains and networks. This stage releases complexing water, which resides in pores.
3. *Polycondensation:* the gel species continue to rearrange and reorganize, forming an increasingly larger network. This results in the three-dimensional aluminosilicate network of the geopolymer binder.

These processes often occur contemporaneously throughout the mixture, rather than in a linear time series. The dense inorganic polymer that forms the geopolymer binder thus provides physicochemical features that may be superior to Portland cement systems, i.e.:

* The interconnected framework gives geopolymers high compressive strengths [13, 14].
* Due to the microstructure of the reaction products, geopolymer and AAB can show good to very good resistance to chemical degradation (sulphate attack, acid attack, seawater attack) [9].
* The geopolymer matrix shows high thermal and fire resistance up to 1000-1200°C [14].
* Geopolymers may exhibit rapid setting without long term deterioration of strength [13].

Of particular interest is the selection of precursor aluminosilicate materials that arise from waste-streams or as by-product pozzolans, which are readily available from existing industries [15]. These include pulverised fuel ash (pfa) and ground granulated blast furnace slag (ggbs). Although ggbs has a relatively high demand from an existing market-base, and an associated relatively high value, perhaps equivalent to that of Portland cement, waste-stream pozzolans such as pfa are not fully recycled into value added products, and excesses are stockpiled or landfilled.

Reaction mechanisms responsible for the creation of an amorphous gel from fly ash are complex and still not fully understood [9]. A wide variability in chemical dosages can be found in the scientific literature related to the activation of pfa systems [16], indicating that optimum proportions of alkali species (hydroxides and silicates) in the activating solution, as well as quantity of alkali per binder mass, still need to be investigated to improve our understanding of the reaction mechanisms. Curing parameters such as curing temperature and stand time, i.e. the time elapsed before the start of high temperature curing, play an important role in the full development of the reaction products. Information available in technical literature is relatively limited, and thus a systematic investigation is still needed for determining optimum curing conditions.

Fly ashes are not a “standard product” as their physical and chemical properties vary considerably not only from source to source but also over time from the same power station [9]. The Department of Energy and Climate Change in its digest of UK energy statistics [17] quoted 14 power stations operational at the end of May 2014 that were using coal or coal mixed (gas/oil) fuel, representing 26% of total electricity generated. Heath et al. [18] reported that pfa production in UK is estimated at about 6∙106 t per annum, half of which is sent to landfill, whereas about 114∙106 t of pfa is available in stockpiles. The potential availability for pfa-based concrete production is therefore significant in the UK, although the suitability of such available ashes has never been investigated in a comprehensive manner.

Pfa-based geopolymers need an external energy source in the form of thermal curing for the reaction to take place. This can be a drawback for the upscaling of the process to the industrial level. On the other hand, Ca-rich slags such as ggbs react at room temperature since their reaction, i.e. the hydration of Ca species and the creation of a calcium-aluminium-silicate-hydrate (C-A-S-H) gel, is different from low-calcium precursors. Moreover, the reaction develops at a very rapid pace, often resulting in a very short initial setting time. The blend of pfa/ggbs for achieving a system reacting at room temperature without rapid setting would suit most concrete applications. Relatively few publications are available in the literature [19-23] and thus further investigations are needed to provide a better insight into the properties and performance of such binary systems.

The work described here aimed to study (a) the effect of curing procedure and activator dosages on the strength development of pfa-based mortar, (b) the influence of physical and chemical properties of 13 pfa sources obtained from 8 UK power stations, (c) the effect of partial substitution with ggbs on the compressive strength development and microstructure of the reacted mortar.

# Materials and Methods

The chemical composition of 13 pfa sources from the UK were assessed by X-ray fluorescence (XRF) at the University of Leicester and these are shown in Table 1. Mineralogical characterisation was carried out by X-ray diffraction (XRD) using a PANalytical X'Pert PRO X-ray diffractometer, equipped with programmable anti scatter slits, rotating sample stage and the X’Celerator Real Time Multiple Strip X’lerator Detector. The incident X-rays were from a Copper Kα X-ray source, running at 40 kV and 40 mA and obtained spectra are shown in Figure 1. Quartz (SiO2) and mullite (3Al2O3∙2SiO2) were the main crystalline components identified. The amorphous content (visible as the broad hump in the background of the XRD pattern centred at 20 – 30° 2θ) was quantified by adding an internal standard (corundum) and performing a Rietveld method-based analysis (results are shown in Table 2). Laser diffraction grain size distribution was used to assess the particle size distribution of all the investigated sources of pfa. Results are shown in Figure 2, where the 13 pfa samples are divided in three classes according to their D50 value (shown in Table 2): low (D50 < 25µm), medium (26 < D50 < 30.5µm), and high (D50 > 30.5µm). Pfa A was the only one used to study the effect of curing procedure and activator dosages.

**Table 1.** Elemental composition (XRF) of the 13 pfa sources investigated.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| PFA source | A | B | C | D | E | F | G | H | I | J | K | L | M |
| SiO2 | 51.52 | 48.90 | 44.58 | 49.28 | 48.48 | 56.40 | 51.32 | 43.38 | 51.33 | 51.05 | 51.23 | 49.90 | 51.76 |
| TiO2 | 0.97 | 0.95 | 0.91 | 0.95 | 1.08 | 0.92 | 1.03 | 0.84 | 0.97 | 1.02 | 0.80 | 1.02 | 0.89 |
| Al2O3 | 23.58 | 23.51 | 21.94 | 23.43 | 24.86 | 22.14 | 24.85 | 23.63 | 24.12 | 25.06 | 18.37 | 27.12 | 24.51 |
| Fe2O3 | 7.11 | 12.36 | 10.59 | 10.61 | 9.49 | 6.63 | 7.24 | 6.56 | 8.51 | 8.60 | 4.55 | 7.38 | 8.58 |
| Mn3O4 | 0.07 | 0.15 | 0.13 | 0.10 | 0.12 | 0.07 | 0.09 | 0.13 | 0.11 | 0.09 | 0.06 | 0.08 | 0.15 |
| MgO | 1.75 | 1.97 | 1.93 | 1.73 | 2.22 | 1.59 | 1.56 | 1.36 | 1.92 | 1.79 | 1.36 | 1.45 | 1.92 |
| CaO | 3.86 | 3.41 | 3.61 | 2.68 | 4.53 | 2.97 | 2.88 | 2.83 | 2.75 | 2.37 | 2.55 | 2.25 | 3.78 |
| Na2O | 0.74 | 0.93 | 0.83 | 1.55 | 0.98 | 0.97 | 0.57 | 0.79 | 0.80 | 0.98 | 0.45 | 0.67 | 0.49 |
| K2O | 2.42 | 3.26 | 2.61 | 3.57 | 2.60 | 1.91 | 1.63 | 1.39 | 3.61 | 3.47 | 1.60 | 3.23 | 2.87 |
| P2O5 | 0.37 | 0.41 | 0.38 | 0.32 | 0.45 | 0.58 | 0.75 | 0.72 | 0.25 | 0.33 | 0.61 | 0.33 | 0.25 |
| SO3 | 0.84 | 0.49 | 0.59 | 0.47 | 0.63 | 0.47 | 0.41 | 0.28 | 0.49 | 0.32 | 0.39 | 0.18 | 0.32 |
| V2O5 | 0.09 | 0.06 | 0.06 | 0.07 | 0.07 | 0.05 | 0.06 | 0.03 | 0.06 | 0.07 | 0.03 | 0.06 | 0.05 |
| Cr2O3 | 0.02 | 0.02 | 0.02 | 0.02 | 0.03 | 0.02 | 0.03 | 0.01 | 0.02 | 0.03 | 0.01 | 0.02 | 0.02 |
| SrO | 0.10 | 0.08 | 0.08 | 0.07 | 0.15 | 0.12 | 0.13 | 0.15 | 0.06 | 0.07 | 0.15 | 0.07 | 0.05 |
| ZrO2 | 0.00 | 0.07 | 0.06 | 0.07 | 0.07 | 0.07 | 0.07 | 0.11 | 0.06 | 0.07 | 0.07 | 0.06 | 0.06 |
| BaO | 0.14 | 0.14 | 0.14 | 0.14 | 0.20 | 0.20 | 0.18 | 0.26 | 0.15 | 0.16 | 0.22 | 0.21 | 0.18 |
| NiO | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.01 | 0.01 | 0.02 | 0.01 | 0.02 | 0.01 |
| CuO | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.01 | 0.02 | 0.00 | 0.01 | 0.02 | 0.00 | 0.02 | 0.01 |
| ZnO | 0.04 | 0.04 | 0.03 | 0.04 | 0.06 | 0.04 | 0.06 | 0.02 | 0.04 | 0.04 | 0.01 | 0.06 | 0.01 |
| PbO | 0.03 | 0.04 | 0.03 | 0.05 | 0.04 | 0.05 | 0.06 | 0.02 | 0.04 | 0.04 | 0.02 | 0.05 | 0.02 |
| LOI (%) | 4.92 | 3.60 | 11.17 | 5.01 | 4.07 | 4.61 | 6.99 | 17.40 | 4.62 | 4.53 | 17.28 | 5.20 | 4.28 |
| Total | 98.61 | 100.43 | 99.73 | 100.20 | 100.17 | 99.84 | 99.95 | 99.92 | 99.93 | 100.13 | 99.77 | 99.38 | 100.21 |

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**Figure 1.** XRD spectra for the 13 investigated pfa sources. Labels A-M equate to labels A-M in Table 1.

**Table 2.** Physical and mechanical properties of the 13 pfa sources investigated.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
|  | A | B | C | D | E | F | G | H | I | J | K | L | M |
| D50 (µm) | 23.8 | 29.2 | 41.6 | 27.3 | 20.2 | 26.5 | 30.5 | 37.3 | 32.9 | 17.3 | 34.7 | 13.2 | 91.2 |
| Am.%1 | 82 | 89 | 86 | 90 | 79 | 76 | 75 | 83 | 86 | 87 | 75 | 79 | 81 |
| % 45 µm2 | 66.9 | 63.2 | 51.0 | 69.0 | 77.6 | 62.4 | 61.0 | 54.9 | 57.3 | 76.4 | 56.4 | 84.8 | 18.9 |
| UCS (MPa)3 | 60.9 (0.0) | 47.5 (1.8) | 27.8 (2.6) | 54.3 (0.5) | 61.4 (2.7) | 47.8 (3.5) | 33.6 (5.7) | 48.0 (4.0) | - | - | - | 60.4 (1.6) | - |

*1Amorphous content from XRD Rietveld method quantification*

*2 Volume percentages passing at 45 µm*

*3 UCS = uniaxial compressive strength. Standard deviation in bracket.*



**Figure 2.** Grain size distribution curves for the 13 investigated pfa sources. Labels A-M equate to those in Fig. 1 and Table 1.

Ground granulated blast furnace slag was supplied by Hanson Ltd. The chemical composition was obtained by XRF analysis and detected oxides were CaO (40.62%), SiO2 (35.82%), Al2O3 (13.00%), MgO (8.00%), Fe2O3 (0.53%), TiO2 (0.53%), MnO (0.51%), SO3 (0.05%), with LOI of 0.66%.

Commercially available alkalis were used. Laboratory grade sodium hydroxide (NaOH) in solid form was dissolved in water at the required dosage, whereas sodium silicate was supplied as a solution with the following composition: 12.8% Na2O, 25.5% SiO2 and 61.7% H2O. The alkali dosage (M+) was defined as the percentage mass ratio of total sodium oxide (Na2O) in the activating solution to the binder. The alkali modulus (AM) was defined as the mass ratio of sodium oxide to silica in the activating solution. M+ was from 7.5% to 15%, whereas AM was from 0.5 to ∞ for the first series of experiments. The 0.5 corresponds to an all sodium silicate solution and ∞ corresponds to an all sodium hydroxide solution. A range of dosages were used to find which values gave the maximum strength. These values were then used in subsequent experiments investigating the influence of physical and chemical properties of different pfa sources. For the assessment of the effect of partial substitution with ggbs on the compressive strength development and microstructure of the reacted mortar, AM was fixed at 1.25, whilst the M+ was varied from 3% to 10%.

The water/solids ratio was defined as the ratio between total mass of water (i.e. added water mass + water mass in the alkali solutions) and the total solid mass (i.e. mass of binder + mass of alkali solids). This ratio was kept constant at 0.37, except when the mixture was found to be of very low consistency after mixing. In such cases, additional water was added until a workable mixture was obtained. This was generally only necessary for 100% ggbs binder.

Mortars were produced with a sand/binder ratio equal to 2.75. A 5 l planetary mixer was used. Binder and sand were placed in the mixer bowl and mixed for one minute. The alkali solution was then added and the material was mixed for a further 9 minutes. A flow table test was then performed according to BS EN 12350-5 [24]. Mixing was continued for a further 5 minutes, giving a total mixing time of 15 minutes. The mortar was then transferred to 50 mm cube moulds and compacted on a vibrating table for 60 seconds. All specimens were then sealed by wrapping in polyethylene film.

In the first series of mixes, four sets of curing conditions were used: (a) 1 hour at 20°C followed by curing at 50°C; (b) 24 hours at 20°C followed by curing at 50°C; (c) 1 hour at 20°C followed by curing at 70°C; (d) 24 hours at 20°C followed by curing at 70°C. One cube from each curing regime was tested in compression after 1, 3 and 7 days curing, with two cubes from each curing regime tested at 28 days. Some of the mixes were replicated to confirm the reliability of the results. A 300 kN capacity compression testing machine was used for all testing with a loading rate of 0.8 kN/s.

In the second series of mixes, two sets of curing conditions were used: (a) 1 hour at 20°C followed by curing at 50°C and (b) 1 hour at 20°C followed by curing at 70°C. Same procedure as in the first series of mixes was used for determining the compressive strength.

In the third series of mixes, two sets of curing conditions were used: (a) 1 hour at 20°C followed by curing at 70°C and (b) curing at 20°C (ambient/room temperature). Two replica cubes from each curing regime were tested in compression at 1, 7 and 28 days.

X-ray diffraction on raw materials and reacted pastes was carried out with Panalytical X’Pert Pro MPD Diffractometer with X’Celerator detector scanned the range of 4-70°2θ, using Cu K-α radiation. Zincite was used as internal standard. HighScore Plus software was used to obtain semi-quantitative mineral weight percentages.

Images for microstructural analysis were acquired with either a Philips XL30 Scanning Electron Microscope equipped with an Oxford Instruments INCA energy dispersive X-ray spectroscopy (EDS) system or a Hitachi TM300 SEM equipped with a Bruker Quantax 70 EDS systems. The EDS systems allow the collection of chemical information for spots and areas in the samples. The samples comprised broken pieces (~5.0 x 5.0 x 2.5 mm) of hardened paste mounted onto aluminium stubs using epoxy adhesive, orienting the fractured surface in a convenient position towards the SEM beam also bearing in mind the location of the EDS detector. Samples were coated with a thin gold-palladium layer to provide a conductive pathway to prevent surface electrical charging.

# Results and discussion

This work investigated: (a) the effect of curing procedure and activator dosages on the strength development of pfa-based mortars; (b) the influence of physical and chemical properties of 13 different pfa sources, from 8 power stations in the UK, on compressive strength development; and (c) the effect of partial replacement of pfa with ggbs on the compressive strength development and microstructure of mortars.

## Effect of curing procedure and activator dosages on strength development

Twenty-four combinations of M+ and AM were investigated for each of the four curing regimes, for a total number of 96 different series of mixes. The effects of stand time (i.e. the time elapsed between the mixing and the start of the curing in oven, during which the process of dissolution and gelation of aluminosilicates species takes place), curing temperature, and activator dosage were investigated by determining the compressive strength development.

### Effect of stand time at 20 °C

In order to assess the effect of the time for dissolution and gelation at room temperature on ultimate strength, half the samples in each mix were left to stand for 1 hour prior to oven curing and half for 24 hours. Figure 3 shows the results for the 28-day tests for the all mixes cured at 70 °C. In general, the difference in strength at 28 days between those initially left to stand for 1 hour and those left for 24 hours is small, with the compressive strengths for most mixes being within 6 MPaof each other. The highest alkali dosage (M+ 15%) gave mixed results, with higher strength for 1 h stand time for high silicate solution dosages (AM = 1.25 and 1.5) and the opposite for low silicate dosages (AM = 2 and ∞). This might be explained by the fact that reactive silicates trigger the nucleation of gel and therefore, when a significant amount is present (lower AM), the gel formation can start immediately and it is enhanced by the high temperature. On the contrary, when the addition of silicates is low (higher AM), the system needs first to achieve the dissolution of Si and Al from the precursor material by the alkali solution at room temperature, whereas the thermal treatment starts the solidification of the gel without allowing enough time for the liquid to complete the dissolution process. Although the difference between no stand time and some stand time can make a difference on strengths by increasing the time for early dissolution and gelation, there appears to be no benefit in leaving mixes for longer than 1 hour between mixing and oven curing. This may be of importance when considering transferring the technology to industry.



**Figure 3.** Compressive strengths at 28 days for 1 h and 24 h of stand time of 100% pfa mortars (70°C curing temperature) for a range of alkali modulus (AM) and alkali dosage (M+). Error bars represent ± standard deviation.

### Effect of curing temperature

Figure 4 shows the effect of curing temperature on the 28-day strengths of all mixes. It can be seen that in all cases, with the exception of those with the lowest AM (0.5), 70°C curing temperatures give significantly increased strengths compared to those cured at 50°C for the same time. The same trend was observed irrespective of the “stand time” duration. Mixes with low alkali modulus were extremely viscous and also tended to expand upon thermal curing, forming a muffin like top. Lower curing temperature leads to mitigation of such expansion and this is reflected in the increased strengths.

### Effect of Alkali Dosage

The alkali dosage is a proxy for the alkali concentration, and describes the mass ratio of sodium oxide (Na2O) in the activating solution to pfa, where the water to total reactive solids ratio is fixed.



**Figure 4.** Compressive strengths at 28 days of 100% pfa mortars for 50°C and 70°C curing temperatures (1 h stand time) for a range of alkali modulus (AM) and alkali dosage (M+). Error bars represent ± standard deviation.

Figure 5 shows the effect of alkali dosage on strength development for the six alkali moduli investigated for curing temperature of 70°C and “stand time” of 1 hour. These were the curing conditions that gave the highest compressive strengths. An increase in alkali dosage (M+) resulted in an increase of the strength up to an M+ of 12.5%. Beyond this ‘optimum’ value, the strengths decreased, which is attributed to saturation of the gel with alkali ions resulting in less free water to be available for speciation of silica and alumina oligomers from the dissolution of pfa.

### Effect of Alkali Modulus

The alkali modulus (AM) is the mass ratio of sodium oxide to silica in the activating solution and it is a proxy for the amount of added silica in the activator solution. As can be seen in Figure 6 (curing temperature of 70 °C and “stand time” of 1 hour), there is an optimum range of values for the alkali modulus, above and below which strengths decrease. With increasing alkali dosage, that ‘sweet spot’ broadens out towards higher alkali modulus. Again, results for M+ = 15% do not follow these trends. In general, alkali moduli between 1 and 1.25 give the highest strengths across the alkali dosages investigated.



**Figure 5.** Effect of alkali dosage (M+) for different alkali modulus (AM) on 28-day strength of 100% pfa mortars (1 hour at 20°C, followed by 70°C curing). Error bars represent ± standard deviation.



**Figure 6.** Effect of alkali modulus (AM) for a range of alkali dosages (M+) on 28-day strength of 100% pfa mortars (1 hour at 20°C, followed by 70°C curing). Error bars represent ± standard deviation.

The drop-off in strength with increasing modulus is likely to be due to the reduced amount of available silica that can participate in the ‘reorganisation-gelation-polymerisation’ steps of the geopolymer formation and thus the development of a denser and more complete and long polymer chain. The reduction in strengths at low modulus (AM=0.5: all sodium silicate) can be attributed to the reduction of the pH in the system when only sodium silicate solution is used. The expansion of the cubes upon thermal curing could also have contributed.

Figure 7 shows a 3-D plot clearly identifying the combined effect of alkali dosage and modulus on compressive strength. The ‘sweet spot’ of the optimum alkali modulus and dosage combinations has a strength maximum of 70 MPa at around an alkali dosage of 12.5% and alkali modulus of 1.25.



**Figure 7.** 3-D plot showing the combined effects of alkali dosage and modulus on the 28-day strength of 100% pfa mortars (1 hour at 20 °C followed by 70 °C)

## Influence of physical and chemical properties of raw materials on compressive strength

A comprehensive characterisation of 13 different fly ashes from coal powered electricity generating stations was carried out in order to investigate the suitability of different pfa sources in the UK. Chemical composition, mineralogical composition, grain size distribution and strengths after alkali activation were investigated.

All the pfa samples conformed to the requirements of the BS EN-450 standard [25] with regards to the SiO2+Al2O3+Fe2O3 (SAF), SO3, CaO, MgO, and P2O5 contents, whereas three pfa sources (C, H, K) had LOI higher than the limits prescribed in [25]. Physical and mechanical properties of nine investigated pfa sources are shown in Table 2. It was not possible to cast cubes for testing for four of the investigated pfa sources because the mixes were either too dry (pfa K and M) or flash setting was experienced (pfa I and J). The former appears to be due to the coarse nature of the pfa coupled with a high LOI. The latter may have been suitable for use in concretes if the alkali activator dosage was reduced. This has not however been investigated further.

Optimum activator dosages were used for the geopolymer mortars, i.e. M+ 12.5% and AM 1.25. Compressive strength measured at 28 days for samples cured both at 50°C and 70°C are shown in Figure 8. The previously identified effect of curing temperature on compressive strength development was confirmed when different pfa sources were investigated, with samples cured at 50°C showing lower strength than samples cured at 70°C.

The reaction rate of pfa depends on several chemical and physical factors [9]. The content of Si and Al, the amorphous phase content and the loss on ignition (LOI) are important chemical factors. Particle size of the precursor materials not only affects the fresh properties (water demand, viscosity, setting time, etc.) but also affects its reactivity. The reduction of the grain size through further milling is known as mechanical activation, and it is commonly adopted for increasing the reactivity of powders [26]. The trend between compressive strength and percentage volume passing 45 µm is shown in Figure 9. The value of 45 µm was chosen according to the EN-450 [25] as indicative size for determining the reactivity of the pfa. The strength data were also plotted versus D50, i.e. the value of the particle diameter at 50% in the cumulative distribution, and a similar trend was observed, see Figure 10. The fineness of the pfa appeared therefore to correlate to the compressive strengths, whilst no relationship was observed linking the compressive strength to amorphous content, LOI, or silicon-aluminium-iron oxides (SAF) content for the investigated ashes.



**Figure 8.** 28-day compressive strength of 9 pfa sources at different curing temperatures. Letters A-L equate to same in Table 1 and Figs. 1-2. Error bars represent ± standard deviation.



**Figure 9.** 28-day compressive strength of 9 pfa sources vs. % volume passing 45 µm sieve. Error bars represent ± standard deviation.



**Figure 10.** 28-day compressive strength of 9 pfa sources vs. D50. Error bars represent ± standard deviation.

It may therefore be concluded that one of the most important factors to consider for achieving high compressive strengths is the grain size of the pfa. It must, however, be emphasised that ten out of thirteen of the tested pfa samples are suitable for use in concretes to partially replace Portland cement and thus conform to EN-450. Attempts to examine fuel ash from lagoons were not successful. It may be that the chemical composition of lagoon ash has been altered over time and it is thus considerably different from all the sources investigated in this programme of work.

## Effect of partial replacement of pfa with ggbs

Mortar mixtures containing only pfa did not develop any significant strength at room temperature. However, the addition of ggbs at any level gave significant strength enhancements even at room temperature. Figure 11 shows the strength of pfa/ggbs based mortars with M+ 7.5% and AM 1.25. There is an almost linear relationship between the amount of ggbs in the binder and the strength at 1, 7 and 28 days where cubes were cured at room temperature (20°C). This seems to suggest that pfa did not contribute much to the strength. There was a significant increase in the strength from ~20MPa to ~50MPa with the addition of only 20% ggbs when cured for 1 day at 70 °C. Higher levels of ggbs cured for 1 day resulted in smaller incremental increases up to ~60MPa at 100% ggbs.



**Figure 11.** Effect of ggbs substitution rate on the strength of sodium silicate/sodium hydroxide-activated binders with M+ 7.5% and AM 1.25. (a) 1 day; (b) 7 days; (c) 28 days. Error bars represent ± standard deviation.

Continued curing at 70°C for 7 and 28 days resulted in strength reductions for mixes with greater than 20% ggbs. This is believed to be due to self-desiccation and resulting drying shrinkage, which is an effect that has previously been observed in alkali-activated slags [7]. In accordance with this effect, the highest strength was achieved for specimens cured at room temperature and not for oven-cured samples. Compressive strength of 80 MPa was obtained at 28 days for the mortar containing 80% slag and cured at room temperature.

Subsequent investigations focussed on the effect of alkali dosage on the strength development of pfa/ggbs blends. The AM was kept constant while the M+ was varied from 3% to 10% for varying levels of pfa replacement with ggbs. Figure 12 shows the 3-D plot of the 28-day compressive strength of specimens cured at room temperature. M+ 10% mixes could not be easily cast for ggbs contents higher than 20% due to quick setting. It can be observed that the increase in ggbs content led to an incremental increase in the compressive strength.

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**Figure 12.** 3-D plot showing the combined effects of pfa/ggbs ratio and M+ variation on the 28-day compressive strength of sodium silicate/sodium hydroxide-activated binders cured at room temperature

## X-Ray Diffraction results for raw materials and reacted pastes

In order to investigate the nature and composition of the reaction products, raw material and reacted pastes were analysed with X-ray diffraction (XRD) technique. The raw materials were activated with the following dosages: M+ 7.5%, AM = 1.25 and w/s ratio 0.37. 100% pfa, 100% ggbs and a 50/50 pfa/ggbs paste were produced. After oven curing, pastes were crushed using distilled water in a McCrone Mill for 5 min and dried at 50°C for 14 hours. An internal standard (zincite) was included for semi-quantitative analysis of the material composition. Diffractograms for unreacted pfa showed approximately 75% of amorphous content, with main crystalline phases as quartz (5.8% in weight), mullite (15.8%), plus minor contents of calcite and maghemite. Ggbs showed an amorphous content of about 88%, with about 10% of crystalline calcite and less than 2% quartz.

Figure 13 shows the compared diffractograms for 100% pfa (A), 100% ggbs (B) and 50/50 pfa/ggbs (C).



**Figure 13.** XRD diffractograms of raw materials and reacted pastes. (a) 100% pfa (b) 100% ggbs (c) 50/50 pfa/ggbs

No new crystalline phase was obtained from the activation of pfa. Quartz and mullite, as expected, did not participate to the reaction, as their mass contents remained unchanged. On the contrary, the activation of ggbs resulted in a decrease in amorphous content from 88% to 78% and in a 50% increase in calcite content from 10.4% to 15.8%, with peaks for calcite clearly higher in the activated ggbs paste diffractogram. It is likely that the calcite in activated ggbs sample is a combination of high and low crystallinity forms, as apparent by the broad base to the 29.4°2θ peak.

The 50/50 pfa/ggbs paste produced results half way between 100% ggbs and 100% pfa pastes. Quartz and mullite contents did not change, whereas the crystalline calcite content increased from 6.2 wt% of the starting material to 10.5 wt% of the paste, which lead to a decrease of the amorphous phase from 82% to 79 wt%.

## Microstructure and chemistry of the pastes

The different mechanical behaviour of pfa and pfa/ggbs mixes discussed above would be expected to correlate in some way with microstructure or reaction products found in the pastes. To test this, SEM analysis was carried out on the three pastes, i.e. 100% ggbs, 100% pfa and 50/50 pfa/ggbs.

### 100% ggbs paste

The basic microstructure of 100% ggbs paste is shown in Figure 14(a) and (b). The paste is made up of unreacted particles of ggbs in a smooth and homogenous looking gel. The paste is traversed by interconnecting curved fractures that locally break up the paste into ~10-20 µm size conchoidal particles. The fractures have morphologies characteristic of desiccation, particularly evident in Figure 14(b). EDS-based chemical analysis of unreacted ggbs and the surrounding gel from the points indicated in Figure 14(a) are summarised in Table 3 and on the ternary diagram in Figure 15. The vertices of the ternary diagram have been chosen to encompass the main chemical compositions of the gels developed in all three pastes. The Na, Ca and Al values in Table 3 have been ‘normalised’ to Si content in each case in order to alleviate the inherent semi-quantitative nature of EDS chemical analysis on rough surfaces. It is worth noting in Figure 15 that the unreacted 100% ggbs particles and the 100% ggbs gel form separate clusters that lie on a line that intersects the Na/Si corner of the plot, where the alkali activator plots (see dashed line in Figure 15). The 100% ggbs gel thus lies about half way along a potential mixing line between the original ggbs composition and the activator composition, suggesting more or less equal contribution of ggbs and activator in the reaction to produce the gel.

****

**Figure 14.** SEM back-scattered electron (A-G) and secondary electron (H) images of geopolymer paste broken surfaces. Numbered spots on A, C, D, E and F are locations of EDS chemical analyses. A & B: paste made with 100% ggbs. C&D: paste made with 100% pfa. E-H: paste made with 50% ggbs and 50% pfa.

**Table 3.** EDS analysis of 100% ggbs paste. Spot numbers refer to Figure 14(a). Analyses with Oxford Instruments INCA.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Spot** | **O** | **Na** | **Mg** | **Al** | **Si** | **K** | **Ca** | **Ti** | **Mn** | **Fe** | **Al/Si** | **Ca/Si** | **Na/Si** | **Description** |
| 1 | 53.42 | 20.99 | 2.00 | 3.22 | 9.96 | 0.15 | 9.95 | 0.13 | 0.17 |  | 0.323 | 0.999 | 2.107 | C-(N)-A-S-H gel |
| 2 | 50.95 | 16.29 | 3.13 | 4.03 | 11.07 | 0.31 | 13.69 | 0.27 | 0.23 |  | 0.405 | 1.237 | 1.472 | C-(N)-A-S-H gel |
| 3 | 46.93 | 20.21 | 1.07 | 3.00 | 11.38 | 0.42 | 16.63 | 0.23 |  |  | 0.301 | 1.461 | 1.776 | C-(N)-A-S-H gel |
| 4 | 49.80 | 19.25 | 1.38 | 3.18 | 11.25 | 0.27 | 14.20 | 0.23 | 0.28 | 0.16 | 0.319 | 1.262 | 1.711 | C-(N)-A-S-H gel |
| 5 | 43.54 | 5.30 | 3.60 | 5.80 | 15.09 | 0.41 | 25.23 | 0.28 | 0.60 |  | 0.582 | 1.672 | 0.351 | Unreacted ggbs particle |
| 6 | 43.57 | 7.08 | 3.65 | 4.94 | 13.72 | 0.53 | 24.72 | 0.60 | 1.05 |  | 0.496 | 1.802 | 0.516 | Unreacted ggbs particle |
| 7 | 36.59 | 3.71 | 3.14 | 5.22 | 15.97 | 0.61 | 33.99 | 0.56 | 0.56 |  | 0.524 | 2.128 | 0.232 | Unreacted ggbs particle |

**Table 4(a).** EDS analysis of 100% pfa paste. Spot numbers refer to Figure 14(c). Analyses with Oxford Instruments INCA.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Spot** | **O** | **Na** | **Mg** | **Al** | **Si** | **K** | **Ca** | **Ti** | **Mn** | **Fe** | **Al/Si** | **Ca/Si** | **Na/Si** | **Description** |
| 1 | 47.27 | 2.78 | 0.76 | 15.02 | 26.5 | 3.58 | 0.94 | 0.78 |  | 2.36 | 1.508 | 0.035 | 0.105 | Unreacted pfa particle |
| 2 | 54.2 | 8.53 | 0.54 | 11.15 | 19.08 | 1.51 | 2.08 | 0.28 |  | 2.62 | 1.119 | 0.109 | 0.447 | N-A-S-H gel |
| 3 | 49.77 | 10.24 | 0.58 | 8.96 | 22.88 | 2.18 | 2.7 | 0.51 |  | 2.17 | 0.900 | 0.118 | 0.448 | N-A-S-H gel |

**Table 4(b).** EDS analysis of 100% pfa paste. Spot numbers refer to Figure 14(d). Analyses with Oxford Instruments INCA.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Spot** | **O** | **Na** | **Mg** | **Al** | **Si** | **K** | **Ca** | **Ti** | **Mn** | **Fe** | **Al/Si** | **Ca/Si** | **Na/Si** | **Description** |
| 1 | 53.09 | 1.21 | 0.89 | 14.18 | 23.50 | 2.89 | 1.34 | 0.44 |  | 2.37 | 1.424 | 0.057 | 0.051 | Unreacted pfa particle |
| 2 | 47.03 | 14.67 | 0.53 | 9.57 | 20.68 | 1.94 | 1.72 | 0.42 | 0.23 | 3.20 | 0.961 | 0.083 | 0.709 | N-A-S crystal |
| 3 | 48.34 | 18.89 | 0.49 | 7.77 | 17.44 | 1.93 | 1.63 | 0.53 |  | 3.03 | 0.780 | 0.093 | 1.083 | N-A-S crystal |
| 4 | 51.39 | 7.02 | 0.76 | 11.05 | 21.23 | 1.93 | 2.11 | 1.18 |  | 3.29 | 1.109 | 0.099 | 0.331 | N-A-S-H gel |

**Table 5(a).** EDS analysis of 50/50 pfa/ggbs paste (1st sample). Spot numbers refer to Figure 14(e). Analyses with Oxford Instruments INCA.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Spot** | **O** | **Na** | **Mg** | **Al** | **Si** | **K** | **Ca** | **Ti** | **Mn** | **Fe** | **Si/Al** | **Ca/Si** | **Na/Si** | **Description** |
| 1 | 38.00 | 0.27 | 4.86 | 6.98 | 18.58 | 0.51 | 29.44 | 0.63 | 0.46 | 0.26 | 0.701 | 1.584 | 0.015 | Unreacted ggbs particle |
| 2 | 29.37 | 0.37 | 4.41 | 7.49 | 19.39 | 0.51 | 37.24 | 0.61 | 0.39 | 0.23 | 0.752 | 1.921 | 0.019 | Unreacted ggbs particle |
| 3 | 45.51 | 1.25 | 1.17 | 18.27 | 24.38 | 2.19 | 1.22 | 0.64 |  | 5.39 | 1.834 | 0.050 | 0.051 | Unreacted pfa particle |
| 4 | 40.95 | 0.47 | 4.53 | 6.95 | 17.02 | 0.47 | 28.54 | 0.40 | 0.32 | 0.37 | 0.698 | 1.677 | 0.027 | Unreacted ggbs particle |
| 5 | 49.12 | 8.41 | 1.21 | 7.04 | 20.92 | 1.28 | 10.44 | 0.39 |  | 1.06 | 0.707 | 0.499 | 0.402 | C-(N)-A-S-H gel |
| 6 | 48.95 | 5.10 | 2.83 | 6.42 | 17.77 | 1.03 | 16.37 | 0.31 | 0.38 | 0.87 | 0.645 | 0.921 | 0.287 | C-(N)-A-S-H gel |
| 7 | 40.81 | 5.41 | 2.20 | 6.96 | 21.86 | 1.55 | 19.17 | 0.44 | 0.49 | 1.12 | 0.699 | 0.877 | 0.247 | C-(N)-A-S-H gel |
| 8 | 39.53 | 6.31 | 1.93 | 8.45 | 23.71 | 1.55 | 15.8 | 0.39 | 0.5 | 1.84 | 0.848 | 0.666 | 0.266 | C-(N)-A-S-H gel |

**Table 5(b).** EDS analysis of 50%/50% pfa/ggbs paste (2nd sample). Spot numbers refer to Figure 14(f). Analyses with Bruker Quantax 70.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Spot** | **O** | **Na** | **Mg** | **Al** | **Si** | **K** | **Ca** | **Ti** | **Mn** | **Fe** | **Si/Al** | **Ca/Si** | **Na/Si** | **Description** |
| 1 | 44.01 | 1.20 | 4.42 | 6.33 | 17.39 | 0.52 | 24.98 | 0.46 | 0.55 | 0.13 | 0.636 | 1.436 | 0.069 | Unreacted ggbs particle |
| 2 | 44.37 | 3.06 | 1.19 | 16.12 | 26.12 | 3.51 | 2.16 | 0.37 |  | 3.10 | 1.618 | 0.083 | 0.117 | Unreacted pfa particle |
| 3 | 51.08 | 7.82 | 1.08 | 6.88 | 18.22 | 1.85 | 12.79 | 0.27 |  |  | 0.691 | 0.702 | 0.429 | C-(N)-A-S-H gel |
| 4 | 49.55 | 7.53 | 0.86 | 7.16 | 18.21 | 2.14 | 12.67 | 0.57 | 0.41 | 0.89 | 0.719 | 0.696 | 0.414 | C-(N)-A-S-H gel |
| 5 | 46.36 | 8.01 | 1.54 | 7.12 | 20.92 | 0.96 | 13.64 | 0.24 |  | 1.20 | 0.715 | 0.652 | 0.383 | C-(N)-A-S-H gel |
| 6 | 49.03 | 8.32 | 1.78 | 7.18 | 20.29 | 2.05 | 10.17 | 0.29 | 0.25 | 0.65 | 0.721 | 0.501 | 0.410 | C-(N)-A-S-H gel |
| 7 | 51.84 | 7.97 | 1.54 | 5.95 | 17.14 | 1.54 | 12.17 |  | 0.89 | 0.97 | 0.597 | 0.710 | 0.465 | C-(N)-A-S-H gel |

### 100% pfa paste

The basic microstructure of 100% pfa paste is shown in Figure 14(c) and (d). The paste is made up of unreacted particles of pfa in a rough and sugary looking granular gel. There are commonly gaps between the gel and the unreacted pfa suggesting weak bonding of gel to unreacted particles. Some long fractures are present, but these have a morphology suggesting they formed during breakage of the sample rather than by desiccation. EDS-based chemical analysis of unreacted pfa and the surrounding gel from the points indicated in Figure 14(c) and (d) are summarised in Table 4(a) and (b) as well as on the ternary diagram in Figure 15. It is clear that the pfa cenospheres are relatively enriched in Al and depleted in Ca and Na relative to Si content. On the other hand, the rough sugary gel is relatively enriched in Na and slightly enriched in Ca relative to the unreacted pfa. In this case, the unreacted pfa and the gel do not lie directly on a potential mixing line between the unreacted pfa and the Na/Si corner of Figure 15. While this may be due to analytical errors resulting from EDS-analysis of rough surfaces, it is also possible that reactions responsible for gel formation preferentially fractionated available Ca in the pfa into the gel.

### 50% pfa and 50% ggbs paste

The basic microstructure of 50% pfa and 50% ggbs paste is shown in Figure 14(e) and (f). The paste is made up of unreacted particles of pfa and ggbs in a smooth isotropic looking gel. The microstructure for 50/50 pfa/ggbs shows unreacted or partially reacted pfa cenospheres and ggbs grains surrounded by an amorphous matrix which was smooth and homogeneous in some places whilst rough and more heterogeneous in others. There are relatively few fractures, which do not look like desiccation fractures. The contact between gel and unreacted ggbs is typically sharp, whereas there is often a gap between the gel and unreacted pfa cenospheres; the large cenosphere in the centre of Figure 14(h) is an exception. EDS-based chemical analysis of unreacted pfa/ggbs and the surrounding gel from the points indicated in Figure 14(e) and (f) are summarised in Table 5(a) and (b) as well as on the ternary diagram in Figure 15. The unreacted ggbs and pfa have compositions similar to those in the 100%ggbs and 100%pfa pastes, respectively (Figure 15). The gel has a composition that lies within a triangle formed by the unreacted pfa and ggbs and the alkali-activator composition indicating all three reactants were involved in the reaction to produce the gel. The position of the gel composition relative to the reactants indicates ggbs was the main contributor to the gel and the alkali activator the least.



**Figure 15.** Triangular plot of Ca, Al, Na compositions normalised to Si content for PFA and GGBS particles, various gels, and some Na-Si crystals formed in one sample. Calculated compositions for the pastes are plotted as six pointed stars. Added NaOH and sodium silicate solution would plot on the Na/Si corner. See text for further explanation and discussion.

The three microstructures are quite different. The 100% pfa gel looks granular and potentially porous with a commonly poor adhesion to unreacted pfa particles. In contrast the 100% ggbs gel is homogeneous and looks denser, but is cut by interconnecting desiccation cracks. Both microstructures look inherently weak. The 50%pfa-50%ggbs gel microstructure is homogeneous and dense looking and lacks the desiccation cracks found in the 100% ggbs gel. The addition of aluminium from the pfa has resulted in a non-granular gel that does not shrink, and is perhaps stronger.

Overall, the main product from a pfa-based system is a sodium-alumina-silicate-hydrate geopolymeric gel (N-A-S-H) with very low Ca/Si ratio, though perhaps a little higher than would be expected from the compositions of pfa and the alkali activator. The gel composition is similar to the calculated bulk composition of the gel ingredients (Figure 15).

In contrast, the high calcium 100% ggbs system produces a calcium-alumina-silicate-hydrate gel (C-A-S-H) with low Al/Si ratio of 0.3-0.4 and a Ca/Si of 1 to 1.5, as has been reported by several researchers [27-29] for alkali activated slag systems. The reaction products are similar to the C-S-H gel in Portland cement systems, where higher Ca/Si ratios are obtained in the range 1.2 to 2.3 [30], and are presumably various forms of calcium silicate hydrates and calcium aluminium hydrates. The gel composition is much more Na-rich than the calculated bulk composition of the gel ingredients and even the unreacted ggbs particles contain significant Na (Figure 15). Re-examination of the same surface about 9 months later revealed a coating of sodium carbonate crystals in rosette forms (Figure 16) indicating an excess of Na in the gel reaction. These rosettes were not visible shortly after the 100% ggbs paste was made (see Figure 14(a) and (b)). This raises the possibility that the higher than expected Na measurements from the surfaces in Figure 14 (a) and (b) reflect initial sub-microscopic formation of sodium carbonate crystals at the sample surface as efflorescence due to evaporation of excess sodium-bearing solute, which as it progressed led to formation of the larger rosette crystals. Thus, we suggest that the 100% ggbs gel at the broken sample surface has progressively incorporated Na from the alkali activator.

The 50% pfa 50% ggbs system produces a C-A-S-H gel with Na/Si in the range 0.25-0.5, Ca/Si at 0.5-0.9 and Si/Al at 0.6-0.85, compositions similar to the calculated bulk composition of the gel ingredients (Figure 15). The gel microstructure looks dense and is uncracked. It has been reported in [31] that alkalis (Na in this case) can be incorporated into a C-A-S-H gel structure in order to balance the charge of tetrahedral Al, therefore it could also be written C-(N)-A-S-H. As suggested in the literature [32], the calcium silicate hydrate gel fills the pores and voids left by the water escaping the “pure” geopolymeric gel (N-A-S-H), resulting in a more compact and dense structure that is responsible for the increase in mechanical strength. The coexistence of N-A-S-H and C-A-S-H gels has been reported in the literature [29, 32-33].



**Figure 16**. Back-scattered electron images of 100% ggbs paste surface ~9 months after images in Figure 14. A shows distribution of sodium carbonate crystals (dark areas) grown on the paste surface (light area). B shows a detail of the rosette structure formed by the sodium carbonate crystals and the characteristic desiccation cracks in the gel.

# Conclusions

Geopolymer and alkali activated binders can offer a possible alternative to Portland cement concrete. Notwithstanding all the research work that has been carried out all over the world, an easy way of selection of raw material and their proportioning to obtain desired properties has remained a trial and error procedure. The following factors affecting the mechanical properties were therefore investigated in order to provide guidance for the selection of raw materials and their proportioning to produce geopolymer concretes of the required properties: (a) the effect of curing procedure and activator dosages on the strength of pfa-based mortar; (b) the influence of physical and chemical properties of different pfa sources; (c) the effect of partial pfa replacement with ggbs.

The conclusions from this work are:

1. Curing temperature has a very significant effect on strength of pfa based geopolymers: specimens cured at 70°C were considerably stronger than specimens cured at 50°C. Stand time was found to be less important and as such an 1 hour stand time before oven curing was considered sufficient.
2. The dosage of activators is very important for not only achieving the required early age properties but also for the effect on compressive strength. A ‘sweet spot’ of the optimum alkali modulus and dosage combinations, i.e. alkali dosage of 12.5% and alkali modulus of 1.25, gave compressive strength of ~70 MPa.
3. Physical and chemical properties of potential pfa sources should be investigated before selecting the most suitable one. Average grain size was found to be one of the important factors affecting the potential compressive strength. Coarse pfa coupled with low amorphous content and high LOI needs to be avoided.
4. Partial pfa replacement with ggbs leads to increases in the compressive strength. Strengths of 80 MPa with only M+ 7.5% and AM 1.25 were obtained. The other benefit from such blends is that curing at room temperature only is sufficient and no elevated curing temperatures are needed.
5. SEM investigation on pastes of 100% pfa, 100% ggbs and 50/50 pfa/ggbs assisted in determining the microstructure, reaction products and their chemical composition. The 100% pfa and 100% ggbs mixes produce distinctly different microstructures, i.e. dominated by coarse granularity and desiccation cracking respectively. The 50/50 pfa/ggbs mix develops a homogeneous, dense gel with little microcracking and good bonding to unreacted ggbs and pfa particles. The 50/50 pfa/ggbs microstructure looks like it should be stronger and it has been proved to be so.
6. A calcium alumina silicate hydrate gel with inclusions of Na+ cations in the structure was found in the samples containing ggbs. This was denser than the sodium alumina silicate hydrate gel (N-A-S-H) found for 100% pfa samples and this could explain the improved compressive strengths.

# Acknowledgements

This research was carried out at the University of Liverpool in the framework of the Carbon Trust Applied Research Grant 0911-0252 “Ultra High Performance Fibre Reinforced Cementless Precast Concrete Products”. The work was then continued at Queen’s University of Belfast with the financial support of the SUSCON project, which has received funding from the European Union Seventh Framework Programme (FP7/2007-2013) under Grant Agreement No. 285463 (Call FP7-2011-NMP ENV-ENERGY-ICT-EeB).

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