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Effect of synthesis parameters on the performance of alkali-activated Nonconformant EN 450 Pulverised Fuel Ash

S. Adu-Amankwah, J.M. Khatib, D.E. Searle, L.Black

Abstract

The fly ash reported in this paper is coarser than conventional pulverised fuel ash (PFA), with loss on ignition (LOI) exceeding 10.8%. Consequently, it is precluded from being used as a supplementary cementitious material (SCM) according to EN 450 and disposed in landfills. Alkali-activation of such PFAs is considered here. Three concentrations of sodium hydroxide (NaOH) were separately blended with water glass at different ratios to modify the silica modulus. Heat of reaction, setting time, compressive strength and drying shrinkage were investigated as a function of activator composition. Specimens were either cured at room temperature or hydrothermally treated at 75 °C for five hours. The results show that by optimizing the activator composition, a binder with a 28 day compressive strength of 25MPa can be synthesised from such PFAs even at room temperature. Among the activator parameters, the alkali content was observed to be most influential.

Keywords: Geopolymer, fly ash, loss on ignition, shrinkage, setting time, EN 450

1.0 Introduction

Alkali-activated binders present a low-carbon alternative to Portland cement [1]. This class of binders are obtained from the reaction between alumina-silicate feedstock and concentrated solutions of alkali hydroxides [2, 3], silicates [4], sulphates [5], carbonates [6] or combinations thereof [7, 8]. An overview of the mechanism responsible for the transformation of the initial constituent materials into a binder has been extensively reported [9, 10]. At higher pH, reactive alumina and silicates dissolve from the feedstock and when speciation equilibrium conditions are attained, the species precipitate into gels. Progressive interconnectivity of precipitated species, reorganization and condensation lead to a solid binder phase.

The parameters which influence the fresh and hardened properties of alkali-activated binders fall into two categories. That is, those inherent within the constituent materials, and the externally applied conditions. With regards to the intrinsic parameters, a variety of activators have been studied [11-14], as have a variety of feedstock materials [13-15]. The proportioning of these materials are system specific [16, 17] and consequently determine the properties of the binder produced [3]. The effect of silica moduli [4, 18, 19], silicate to alumina [20-22] and water to silicate ratios have all been explored. However, mechanical performance is usually related to the silica modulus [23, 24]. Recent studies have suggested that the amorphous silica to alumina ratio (SiO₂/Al₂O₃) also plays a significant role on the early age properties [25]. The optimum SiO₂/Al₂O₃ ratio however depends on the calcium content of the feedstock [26, 27]. It therefore follows that a generalised mix formulation in these binders does not exist, thus necessitating the need to optimize the activator composition for every feedstock.

A range of alumina-silicate source materials for the synthesis of this class of binders have been explored in the literature [28-31]. Most of the explored feedstock however requires thermal or mechanical processing in order to obtain reasonable performance [32-37]. As a result, pulverised fuel ash (PFA), an industrial by-product which can be used directly will remain a popular and rather cheaper option. By virtue of being an industrial waste, the properties of PFA are dependent on coal source [25], pulverisation and combustion conditions [38, 39] which are often tailored to maximize energy output rather than the quality of the ash. Technology and control of environmental pollution also dictates the ash collection method which also impacts on the quality of the PFA [40]. This leads to wide compositional variability, even from a single power stations [41, 42].

The variability of PFAs even from the same source restricts the use of some PFAs as SCMs. For example, the European norm EN 450-1 imposes compositional and physical requirements for PFA as SCMs; unburnt carbon content below 9%, as well as a maximum content of 40% coarser than 45 microns [43]. These non-conformant PFAs end up as landfill wastes at a cost to power generators. Previously reported alkali-activated PFAs offer potential technical advantages. However, the PFA feedstock mostly conforms to the EN 450 criteria [3, 44-48] thus competing with those used as SCMs in composite cements. The suitability of Non conformant PFA for geopolymer synthesis has not been studied previously and constitutes the objective of this paper.

2.0 Experimental Details

2.1 Materials

Unconditioned PFA obtained directly from electrostatic precipitating hubs from a UK Power station was used for this study. The X-ray fluorescence (XRF) composition and the particle size distribution obtained through Malvern Mastersizer are shown in Table 1 and Figure 1 respectively.

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Oxide	SiO ₂	Al ₂ O ₃	Fe2O	CaO	MgO	Na ₂ O	K2O	TiO ₂	SO ₂	LOI
%	51.42	24.33	13.10	4.06	2.16	0.85	2.45	1.15	0.52	10.78

Table 1 XRF composition of the PFA under investigation

Iso-propanol was used as dispersant for the PSD measurement and the Fraunhofer method was implemented for data evaluation. The LOI was separately determined according to the method in EN 196-2 hence the sum of the oxide compositions including the LOI exceeding 100%.

Activators for the study were formulated from water glass solution and laboratory grade sodium hydroxide pellets (98% purity). The water glass solution composed of 8.5 ± 0.3 % Na₂O and 27.8\pm0.5% SiO₂. The viscosity and specific gravity were 70-120MPa.s and 1.385g/cm³ respectively. Three concentrations of sodium hydroxide solutions (8, 15 and 20M) were added at different levels to modify the silica modulus

[4]. The activator composition and calculated molar ratios are shown in Table 2. The choice of the investigated activator composition range was based on the varied optimal ranges reported in the literature [49].

2.2 Sample preparation and testing

Isothermal conduction calorimetry was conducted on an 8-channel TAM Air calorimeter. 6.0g of PFA and 3.0g of the activating solutions were weighed and kept in the calorimeter until a stable baseline was attained. The activating solution was subsequently injected and then mixed in situ.

Setting time was measured on paste samples according to BS EN 196-3 using the Vicat apparatus. The entire test setup was submerged under water as prescribed in the norm. The reported setting times are average of three measurements. Mixing procedure for setting time, shrinkage and compressive strength testing were based on the recommendations of BS EN 196-1. The activating solution to feedstock ratio was maintained at 0.5. This was established from the consistency tests during the scoping studies.

Compressive strength and drying shrinkage were measured on mortar samples in The mortar specimens were prepared with 1:1:0.5 replicates of three. PFA/sand/activator ratios. The mix ratio was adopted in order to maintain the activating solution/PFA ratio at the same level as that which was used for the setting time and calorimetry measurements. The specimens for strength test were 50mm cubes. Shrinkage test was based on BS ISO 1920-8:2009 but using 40x40x160mm mortar prisms and demountable mechanical (i.e. demec) gauge studs. Measurements commenced 1 day after the relevant curing method. The demec points were attached on all four longitudinal sides of the test specimens using the polyester-based adhesive, chemical metal. Autogenous shrinkage (Auto) was taken as the shrinkage on samples sealed with flash band and monitored from 1 day. Two curing regimes were assessed; continuous curing at room temperature (RT) and hydro-thermally treated at 75°C for five hours (75), as performed elsewhere [50]. The hydro-thermal curing was achieved by introducing the moulds in a pan which was half-filled with water and covered with an aluminium foil.

Designation	А	В	С	D	Е	F	G	Н	J	к	L	М	Ν	Ρ	Q
NaOH conc Group1 waterglass/8M					Group2 waterglass/15M					Group3 waterglass/20M					
SiO ₂ /Na ₂ O	0.5	1.1	1.7	2.6	3.1	0.4	0.8	1.4	2.3	3.0	0.3	0.7	1.2	2.2	2.9
Molar H ₂ O	5.4	5.2	5.0	4.9	7.5	4.7	4.7	4.8	4.8	7.4	4.4	4.5	4.6	4.7	6.3
H ₂ O/Na ₂ O	21	23	27	31	52	12	16	20	26	50	10	13	17	25	41

Table 2 Activator composition and computed parameters

H ₂ O/SiO ₂	39	23	16	12	17	34	20	15	12	17	31	19	14	11	14
w/solids	0.3	0.3	0.3	0.2	0.4	0.2	0.2	0.2	0.2	0.4	0.2	0.2	0.2	0.2	0.3

Note: Entries rounded to 1 decimal place.



Figure 1 Particle size distribution of the PFA under investigation

3.0 Results and discussion

3.1 Reaction kinetics

The data presented here is limited to 20°C because it was not possible to perform hydrothermal curing in a calorimeter. Previous studies also converge on enhanced kinetics of reaction at elevated temperatures [3, 51, 52]. The objective here is therefore to clarify the influence of alkali concentration and dosage on the reactivity and kinetics at ambient temperature.

The kinetics of early stage reactions as a function of activator composition are shown in Figures 2 (a-c). In all formulations, the heat flow was characterized by a single exothermic peak which occurred within 20 minutes after mixing. A short induction stage can be seen in the 8M NaOH mixes (Figure 2a); but no appreciable induction was observed in the 15M and 20M mixes. The end of the induction period which was about 3 minutes in the present study marked the onset of the dissolution of silicate and aluminate species [24] from the PFA. This being distinct in the 8M NaOH activated mixes can be explained by the differences in the aluminosilicate dissolution rate [53, 54]. The 8M NaOH activated mixes contained more water but lower hydroxyl ions compared to the 15M and 20M NaOH blends. Consequently, the increased PFA/hydroxyl ion ratio retards dissolution. This is consistent with the data from leaching experiments reported elsewhere [55].

At a given NaOH molarity, an inverse relationship was noticed between the silicate dosage and the rate of reaction. This effect was less distinct in the 8M NaOH mixes except at silica moduli of 2.6 and 3.1. Comparison between Figures 2a and 2b depicts the significance of the alkali concentration which was used to modify the silica modulus of water glass. The rate of reaction was accelerated with increasing alkali concentration. However, this seems to reach a maximum at 15M NaOH. Beyond this, differences between 15 and 20M NaOH were not visible. This may be attributed to comparable dissolution and polymerization rates at a threshold hydroxyl concentration [56]. Beyond this threshold, further increase in hydroxyl concentration has little or no influence on the overall reaction rate.





Figure 2 Effect of alkali concentration and activator composition on the heat flow of alkali-activated RoS PFA: (a) 8M NaOH blends; (b) 15M NaOH blends; (c) 20M NaOH blends. Refer to Table 2 for designations

In the synthesis of alkali-activated binders, the activator is often specified in terms of the silica moduli [24, 44, 57]. The general trend of improved rate of reaction with decreasing silica modulus is consistent with the literature [24]. However, the heat flow data presented above have shown that, slight modifications in the silica modulus can influence the reaction kinetics significantly. The mechanisms underlying these observations seem to relate to the molarity of the alkali solution rather than the silica modulus. The heat flow data is consistent with the previously espoused geopolymerisation mechanism [9, 44]. The dissolution, polymerization and condensation processes are coupled. However, dissolution is exothermic and accounts for the single heat flow peak. Subsequent reactions seem to produce minimal net heat of reaction after the first 40 minutes. Some authors [24, 56] separated the reaction rate peak into acceleration and deceleration stages which correspond to dissolution and precipitation of reaction products respectively. The maximum heat evolved for each of the three groups was plotted against the silica modulus and shown in Figure 3. This is only used as a proxy to indicate the extent of dissolution in the systems investigated and fully recognizes possible concurrent dissolution and precipitation of alumina-silicate gels [9]. The trends are similar irrespective of the alkali concentration. The maximum heat decreased with increasing silica modulus. At any given moduli, the maximum heat was similar at 15 and 20M but lower at 8M.



Figure 3 Effect of the silica modulus on the maximum heat evolved in activated RoS PFA

At a given molarity, increasing the dosage of the original waterglass solution reduces alkalis and the hydroxyl content. Consequently dissolution of aluminate and silicate species from the precursor is restricted. The lower evolved heat in the 8M NaOH blended binders can be explained by the suppressed dissolution and gelation at lower alkalinity [29, 55]. Recent studies reported elsewhere [58] have confirmed that persistence of aluminate species in solution inhibits further dissolution from the feedstock.

In the aqueous solution, hydrolysed silicates may be protonated and the nature may vary from monomers to oligomers [59]. The prevalence of oligomeric silicates result in gels which constitute the building blocks for the geopolymer structure. In the interlinked gels, tetrahedrally coordinated [Al(OH)₄]]⁻ substitutes for silicates as polymerization increases. Suppressed dissolution leads to lower fraction of hydroxyl aluminates. Additionally, higher molarity increases the amount of alkalis available to balance the negative charge arising from the substitution of alumina for silicates in the evolving N-A-S-H structure [54].

3.2 Setting time

Final setting times of the investigated mixes as a function of activator composition and total w/solid ratios as measured by the Vicat method are presented in Figures 4 (a-d). Setting time, which is indicative of precipitation of network forming products, is strongly influenced by the composition of the activating solution. The strongest correlation was observed with the silica moduli and the water to alkali ratios while the worst was observed from the water to silica ratio. According to the data, increasing the silica moduli or the water to alkali ratio of the activating solution delayed setting. The accelerating effect of higher alkali concentration on the rate of reaction and consequently quicker setting has been reported elsewhere [60]. Flash setting was observed in the mixes with SiO₂/Na₂O below one. The overall water deficiency at these moduli (SiO₂/Na₂O <1) (see Fig 2-d) potentially resulted in accelerated gelation arising from the pairing of dissolved ionic species in close proximity which then leads to rapid setting. Final setting time was over 6 hours at silica modulus of approximately 3. This may also be attributed to the lower alkali content retarding the dissolution from the feedstock [24, 55].



Figure 4 Initial times of RoS PFA as function of activator parameters: Effect of (a) silica modulus; (b) molar water/ alkali (c) molar water/ silica (d) water/solid ratios

Moreover, at higher silica moduli, there is the thermodynamic tendency for the silicate oligomers to polymerize at the expense of incorporating AI species [14]. This however takes place at a slower rate compared to silicate-alumina [61]. The setting times agree with the heat of reaction data presented in Figure 2 and are consistent with the trends reported in [24]. Despite the importance of alkalis to silicate dissolution, it is worth noting that excess alkalis in solution can also impede the

polymerization of the gels [62]. At SiO₂/Na₂O < 1, the Si/Al ratio of the aqueous solution is low [14] and consequently decreased alumina-silicate oligomers as already explained. Without sufficiently dissolved aluminate species, the polymeric reaction following speciation equilibrium is rather between silicates. Excess alkalinity as in the 20M NaOH blends increases the OH⁻ ions and eventually the water content. These hinder polymerization [63]. However, lower oligomeric alumina-silicates in the gel phase can also be beneficial for the subsequent polymerization reactions [14].

It is important to point out that the Vicat method which measures the penetration resistance of a single needle attached to a 300g plunger is not a direct measure of the evolving sodium-alumina-silicate microstructure. Despite ultrasonic measurement providing reliable assessment of the microstructural evolution [64], the Vicat test is regularly used to characterize the percolation and network formation in cementitious materials[24, 65].

3.3 Compressive strength evolution

Table 3 shows the effects of curing regime and activator composition on compressive strength after 1 and 28 days. Alongside compressive strength, the cumulative heat after 1 day is presented for each group. The impact of hydro-thermal curing on early age strength development was obvious. Compressive strength was higher following hydrothermal treatment compared to room temperature curing. The magnitude of compressive strength however depended strongly on the activator composition.

Mix ID	Ms	H2O/Na2O	Compressive strength, MPa @ 20°C		Compressive strength, Hydrotherma	Cum. Heat @ 1D, J/g of paste	
			1D	28D	1D	28D	
А	0.54	21.37	1.43	4.80	2.91	11.67	13.48
В	1.05	23.77	5.63	14.83	16.31	23.24	15.36
С	1.74	27.02	7.10	12.59	10.82	20.33	16.60
D	2.60	31.12	3.75	5.74	4.67	6.00	9.58
E	3.09	52.10	1.35	2.55	2.10	5.72	9.75
F	0.38	12.61	1.70	11.03	3.24	12.79	19.73
G	0.76	15.62	5.90	16.24	12.99	18.68	21.24
Н	1.37	20.07	11.39	25.53	19.49	35.65	32.91

Table 3 Effects of silica modulus (Ms) and H_2O/Na_2O on compressive strength following curing at room temperature and hydrothermal curing at 75 °C.

J	2.30	26.94	3.79	9.59	4.26	14.92	13.52
К	2.96	49.38	1.81	2.10	1.25	3.73	10.50
L	0.32	10.11	2.65	7.17	5.84	13.11	19.01
М	0.67	12.91	4.34	14.13	6.31	15.68	23.49
Ν	1.23	17.49	8.12	17.30	17.10	29.10	42.98
Р	2.18	25.11	4.54	13.77	4.20	14.26	12.75
Q	2.89	41.19	1.01	1.74	1.21	3.04	10.88

Note: The standard deviation the data points in the table ranged between 0.3 to 0.7MPa.

In the samples cured at room temperature, strength was lowest in the samples that were activated with 8M NaOH blended solutions. Modifying the silica modulus with 20M NaOH did not necessarily improve strength. Slightly lower strength was observed on the 20M NaOH samples which were cured at room temperature. With respect to the silica modulus, it is apparent that, irrespective of curing regime and molarity of sodium hydroxide, an optimum existed. The highest compressive strength was obtained from the moduli range 1.2 - 1.7 and water/alkali ratio of 20. The latter depends more on the alkali content than the initial water or silicate content.

These are consistent with the literature [3, 66]. This study did not reveal any distinct relationship between the H₂O/Na₂O and strength development. Plausible reason is the reaction mechanism [9] in these binders. Here, water which is consumed in the dissolution process reforms during gelation and rearrangement; silicates on the other hand only play a transitional role in the polymerization process [3, 4]. The silicates play a vital role only when dissolution of species from the feedstock is underway. The dissolved silicates in the activator then potentially act as nucleation sites for growth of polymeric chains similar to the effect of seeding on C-S-H evolution in conventional clinker based systems [67]. The nature of the resulting sodium alumina-silicate hydrate (N-A-S-H) varies according to temperature [50] and dissolved Si/Al ratios [63]among other factors.

As noticed from the heat flow data above (see Figure 2), the alkali content strongly influences the kinetics of reaction in these binders and consequently the compressive strength. A threshold alkali concentration relative to dissolved aluminates has however been reported elsewhere [60]. Silica modulus below 1.0 presents highly alkaline systems which leads to rapid super-saturation of dissolved species. This was also manifested in the setting times (Figure 4a). The excess silicates lead to coagulation of the activating solution which inhibits dissolution and subsequent polymerization.

3.4 Drying shrinkage

Figure 4 shows typical total shrinkage and shrinkage due to progressive reactions (measurement commencing after 1 day and subsequently referred to as autogenous shrinkage, denoted 'Auto') for the two curing regimes investigated; hydrothermal curing at 75°C (denoted 75) and continuously at room temperature (denoted RT) for the PFA activated with Silica modulus of 0.8/15M. The data shows that the curing regime strongly influences total drying and autogenous shrinkage. As would be expected, water is evaporated during the hydrothermal curing. The shrinkage associated with the latter was however not taken into consideration. Following the hydrothermal curing, the sample has less water and better developed microstructure. Conversely, the room temperature cured samples reacted slowly with more water available to be removed by drying. The trend between the two curing regimes was consistent irrespective of the activator composition. This implies that, reaction persists in the room temperature cured specimens over the test duration while that of the hydrothermally cured samples were marginal. Higher drying shrinkage in the samples cured at room temperature was expected due to slower rate of reaction and availability of moisture to dissolve CO₂, thus amplifying shrinkage due to carbonation; and also evaporation of water. Carbonation related shrinkage is common in this type of binders and often manifest as efflorescence especially in ambient temperature cured samples [68]. Increasing NaOH concentration in the activator would be expected to increase shrinkage due to leaching of alkalis to form carbonates. The magnitude of this however depends on the distribution and mobility of sodium ions from the N-A-S-H gels.

The effects of activator composition on the net drying shrinkage of the investigated mixes are shown in Figure 5. The trends were similar in the hydrothermally cured samples hence not duplicated. The results show higher shrinkage with increasing silica moduli irrespective of the NaOH concentration used to modify the activating solution. The magnitude of drying shrinkage however showed strong dependence on the alkali concentration as was expected. For example, slight changes in the modulus resulting from using 15M rather than 8M NaOH doubled the drying shrinkage magnitude. However, for approximately similar modulus, the 20M NaOH showed lesser shrinkage compared to the 15M blends. In summary, highest drying shrinkage was noticed when the activator modulus was modified with 15M NaOH and lowest at the 8M NaOH. The shrinkage in the 20M NaOH blends was however higher than the 8M NaOH blends at all times. This means that shrinkage arising from leaching of alkalis is small compared to that due to moisture loss.







Figure 5 Effect of activator composition on the drying shrinkage of room temperature cured specimens. Solid lines: 8M NaOH; dash lines: 15M NaOH; dots= 20M NaOH blends

The observations seem to relate closely to the moisture in the specimen [69]. Considering that the water to alkali ratio increased with increasing silica modulus, higher proportion of water would be available to evaporate at high silica modulus. With regards to alkali concentration, increasing the molarity implied higher solute to water ratio as shown in Table 2; however the shrinkage did not follow this trend. The observations can be attributed to the extent of reaction in the mixes. Given that the hydrolysis of alumina-silicates consumes water but water reforms during condensation [9, 12], higher shrinkage was expected in the samples with greater reaction. This is also consistent with the compressive strength data presented above.

4.0 Conclusions

The effect of activator composition on the reaction kinetics and performance of activated high LOI PFA have been investigated. It has been shown that, by optimizing the activating solution, a binder with up to 25MPa can be produced from EN 450 non-conformant PFA at room temperature. Drying shrinkage was considerably higher than conventional cements. Regarding the activating solution parameters, the silica moduli and water/alkali ratios significantly influenced reaction kinetics, setting time, strength and dimensional stability. Reactivity improved with increasing alkali concentration but decreased at high silicate content of the activating solution. Alkali activation therefore offers the possibility of making good use PFAs which would otherwise be disposed in landfills at a cost and attendant environmental problems.

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