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Alkali-activated Fly Ash Manufactured with Multi-stage Microwave Curing

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ABSTRACT

Alkali-activated fly ash (AAFA) manufactured with conventional thermal curing (85°C) shows comparative mechanical properties to Portland cement (PC) based system, demonstrating its potential as a future low-carbon and environmentally friendly alternative cementitious material. However, the environmental benefit which could have been achieved from the low carbon nature of AAFA is often offset by the higher energy consumption from thermal curing. Since microwave heating is usually considered as a low-energy and low-carbon heating method, some pilot attempts have been made in the literature to cure AAFA with domestic microwave ovens. However, due to the lack of control on the temperature and humidity in the domestic microwave ovens, splashing of fresh AAFA and microcracks of hardened AAFA are two main reasons causing the unsatisfactory performance of microwave cured AAFA.

In this study, a custom-made microwave oven equipped with proportional-integral-derivative (PID) control was employed to cure AAFA samples. The microwave power was automatically regulated through the real-time temperature feedback from the fibre Bragg grating (FBG) sensor embedded in the AAFA sample. The microwave curing regime consisted of four curing stages at four different temperatures, namely 65°C, 85°C, 105°C and 125°C. The similar temperature profile was applied on the AAFA cured in a conventional thermal oven. Compressive strength, reaction products and microstructure of AAFA samples with two curing methods were examined by the analytical techniques of XRD, FTIR, NMR, MIP and SEM respectively. The results showed that AAFA under microwave curing exhibited higher compressive strength with less energy consumption than that under the conventional thermal curing. The microwave curing favoured the formation of Al-rich N-A-S-H gel in AAFA and a more compacted microstructure.

INTRODUCTION

Alkali-activated fly ash (AAFA), consisting of fly ash and alkaline activators, shows equivalent cementitious properties to Portland cement (PC). However, different from PC system, the main reaction product of AAFA is three dimensional amorphous sodium aluminosilicate hydrate (N-A-S-H gel) along with a small quantity of zeolites (Palomo et al., 1999). Compared to PC system, it has been claimed that AAFA produces lower CO₂ emission, consumes less energy and possesses superior durability (Bakharev, 2005a, Fernandez-Jimenez et al., 2006a). However, the strength development of AAFA is very slow under room temperature and thermal curing is essential for the initiation of the chemical reaction and then subsequent strength gain (Puertas et al., 2003, Criado et al., 2005). Currently, the prevailing curing method for AAFA is thermal curing at 85°C, which normally takes more than 8 hours to obtain an early strength of above 20 MPa (Criado et al., 2005). The main drawback of this thermal curing is its high energy consumption, offsetting the environmental benefits which could be obtained from AAFA itself.

Compared with conventional thermal heating, such as thermal oven, it has been generally agreed that microwave is a low energy heating source. The dominant heating mechanism of conventional thermal curing is thermal conduction from the surface to the core of the samples, which is slow and may generate thermal gradient inside the sample (Holman, 2002). Consequently, it can potentially lead to thermal cracks, affecting the strength development. Microwave curing is different from conventional thermal curing. Under the alternating electromagnetic field of microwave, dipolar molecules of dielectric materials can vibrate and, thus, generate frictions between the molecules which then convert microwave energy into heat energy instantly and volumetrically (Metaxas; and Meredith, 1983). As all the components in concrete, such as water, cement and aggregate are dielectric materials, concrete itself can be heated by microwave effectively (Ong and Akbarnezhad, 2015).

Some pilot research have been reported in the literature on using microwave as an alternative heating source to cure cement or concrete products (Wu et al., 1987, Leung and Pheeraphan, 1997). Their results indicated that microwave is a potential low energy technique for manufacturing concrete and both its curing duration and energy consumption can be reduced dramatically. However, the research on using microwave to cure AAFA is still scarce. Somaratna *et al.* (2010) are possibly the only pioneers to cure AAFA mortar in a domestic microwave oven. They found that the compressive strength of AAFA could reach 58 MPa after 120 minutes microwave curing, while only 38 MPa could be obtained after 48 hours thermal curing at 75°C in a thermal oven. Whilst all these previous research have indicated that microwave is a potential low-energy curing technique, one of the main issues facing the previous studies is the lack of an appropriate curing regime and accurate temperature and humidity control during the microwave curing process due to the employment of domestic microwave oven in their studies, which have caused splashing, overheating and thermal cracks inside the samples (Wu et al., 1987). Different from the previous research, a custom-made microwave oven with multi-stage microwave curing regime was employed in this study to cure AAFA. The effects of the proposed multi-stage microwave curing regime on the compressive strength, energy consumption, reaction products and microstructure of AAFA are reported and also compared with those of the AAFA manufactured in a conventional thermal oven in this paper.

EXPERIMENTAL

Materials and custom-made microwave oven

Class F fly ash was used in this study. The chemical composition of fly ash is given in Table 1. Figure 1 shows the SEM image of fly ash particles. The mean particle size of the fly ash is 17.93 µm and the percentage of particles smaller than 45 µm is 88.55%. Sodium hydroxide (NaOH) of industrial grade with

a purity of 95% supplied by ReAgent, UK was used as an activator.

Table 1. Chemical composition of fly ash (wt/%)

SiO ₂	Al ₂ O ₃	Fe ₂ O ₃	CaO	Na ₂ O	MgO	K ₂ O	SO ₃	P ₂ O ₅	TiO ₂	Other	LOI
49.29	30.55	5.55	5.96	0.74	0.81	1.38	0.63	0.21	1.08	0.33	1.87

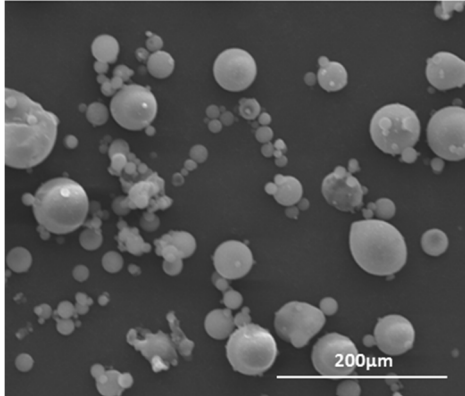


Fig. 1. SEM image of Class F fly ash

The custom-made microwave oven was manufactured by Industrial Microwave System Ltd, UK. The microwave power can be varied from 0 to 1000 Watts continuously. Equipped with fibre Bragg grating (FBG) sensor, the temperature changes inside the sample during the microwave curing process can be recorded and the temperature feedback is then used to regulate the microwave power under proportional-integral-derivative (PID) control.

Preparation of the sample

8M NaOH solution was used in this work to activate fly ash and the NaOH solution to fly ash ratio was 0.33 by mass. Immediately after mixing, the AAFA mixtures were cast into tailored-made plastic moulds, and the fresh AAFA paste samples (25mm×25mm×25mm) were then subjected to different accelerated curing, namely, conventional thermal curing and multi-stage microwave curing. The energy consumption during both curing processes was obtained from the power meters attached to each curing equipment.

The details of each accelerated curing are given below:

- Multi-stage microwave curing (MW): The curing process consisted of four stages (RH>90%) and lasted for different durations, in which the curing temperatures at each stage were 65°C (duration at 65°C, t₁: 35 minutes), 85°C (duration at 85°C, t₂: 35 minutes), 105°C (duration at 105°C, t₃: 35 minutes) and 125°C (duration at 125°C, t₄: 85 minutes) respectively. The curing regime is shown in Figure 2.
- Conventional thermal curing (OV): A similar temperature and curing duration profile as employed in the microwave curing was applied in the conventional thermal curing as a control in the current study;

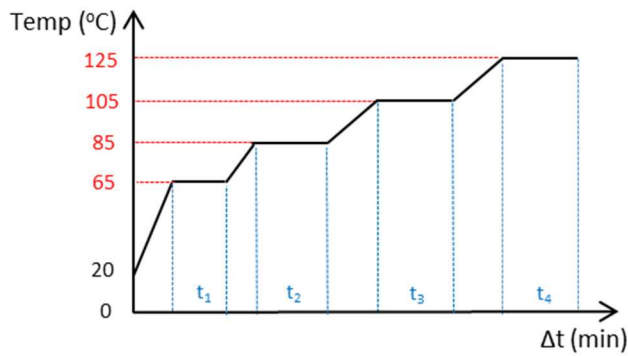


Fig. 1. Multi-stage microwave curing regime for manufacturing AAFA

Tests

The compressive strength of AAFA samples was tested 2 hours after the accelerated curing when the samples were cooled down to room temperature. Reaction products were characterised by means of X-ray Diffraction (XRD), Fourier Transform Infrared spectroscopy (FTIR) and Nuclear Magnetic Resonance spectroscopy (NMR). The microstructure of hardened AAFA was studied by Mercury Intrusion Porosimetry (MIP) and Scanning Electron Microscopy (SEM).

RESULTS

Compressive strength and energy consumption

The compressive strength gained after the accelerated curing as well as the energy consumption incurred during the two curing methods are shown in Table 2. With 4.5 hours multi-stage curing, the MW AAFA obtained higher strength than the OV AAFA, i.e. 35.66 MPa as compared to 28.10 MPa, which is around 27% increase. More importantly, it can be seen that the energy consumption per strength gained (kJ/MPa) is only 191.81 KJ/MPa for the MW AAFA whereas this is 330.92 KJ/MPa for the OV AAFA, which indicates that a 42% reduction in the energy consumption has been achieved from the multi-stage microwave curing compared to the conventional thermal curing.

Table 2. Compressive strength and energy consumption of AAFA with two curing methods

Sample	Curing method	Duration (hour)	Total energy consumption (kJ)	Compressive strength (MPa)	Energy consumption (kJ/MPa)
OV AAFA	Electric oven	4.5	9299	28.10	330.92
MW AAFA	Microwave oven	4.5	6840	35.66	191.81

XRD

Figure 3 presents the XRD spectra of the raw fly ash and the two AAFA samples. The hump appeared in the raw fly ash represents the amorphous phase (as indicated at 17-33° 2θ). The peaks of crystalline phases, such as quartz, mullite and hematite were also detected which is similar to those reported in the literature (Criado et al., 2007a, Kovalchuk et al., 2007). After alkali activation, the area below the hump representing the amorphous phase decreased in both AAFA samples, indicating that the amorphous phase in the raw fly ash was dissolved and took part in the alkali activation (Palomo et al., 1999, Criado et al., 2007a). In addition, two new zeolitic phases were identified in both OV AAFA and MW AAFA. One new zeolitic

phase, as indicated by the new peaks appeared at 13.8° and $24^\circ 2\theta$ were also attributed to hydroxysodalite. The other new zeolitic phase identified at $34.2^\circ 2\theta$ was assigned to chabazite-Na. These results were in good agreement with those findings reported in the literature by other researchers (Criado et al., 2007a, Bakharev, 2005b).

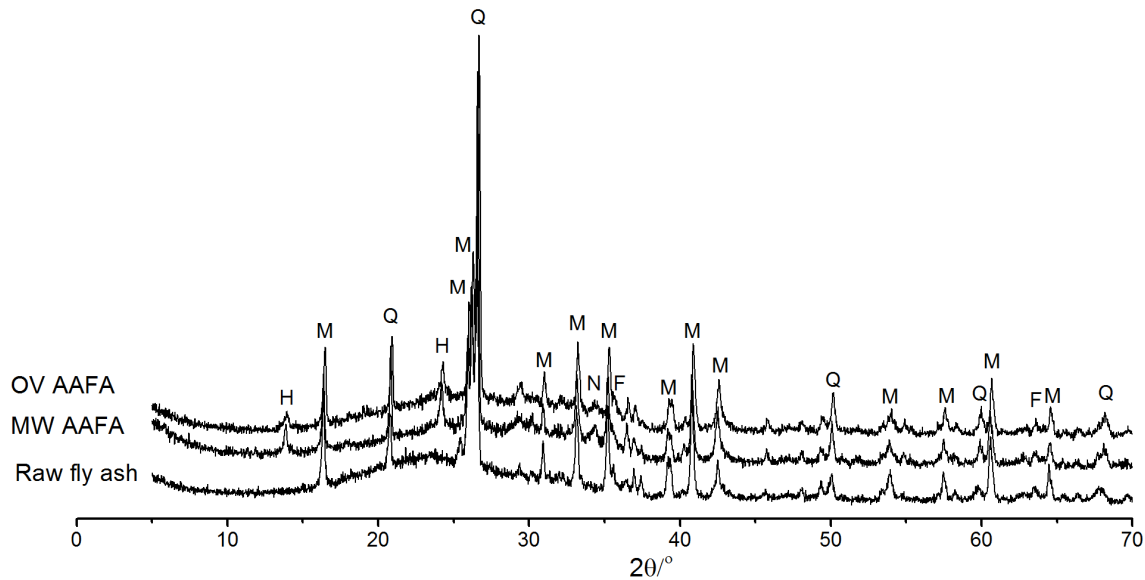


Fig. 3. XRD spectra of raw fly ash and AAFA samples
Q-quartz, M-mullite, F-hematite, H-hydroxysodalite, N-chabazite-Na

FTIR

Figure 4 depicts the FTIR spectra of the raw fly ash and the two AAFA samples. The presence of quartz and mullite in the raw fly ash was responsible for the bands appeared at 791 cm^{-1} and 548 cm^{-1} respectively. In addition, a wide band associated with T-O (T=Al, Si) asymmetric stretching vibrations from the amorphous phase can be observed at 1037 cm^{-1} in the raw fly ash. After alkali activation, the T-O band became sharper and shifted towards lower frequencies at 978 cm^{-1} in OV AAFA and 974 cm^{-1} in MW AAFA respectively, which is considered as the fingerprint of the formation of N-A-S-H gel in AAFA (Fernandez-Jimenez and Palomo, 2005, Criado et al., 2007b). It is interesting to find that the T-O band in the MW AAFA shifted towards even lower frequency than that in the OV AAFA. As the lower frequency of the T-O band in the AAFA indicates more Al tetrahedra is positioned in the N-A-S-H gel (Fernandez-Jimenez and Palomo, 2005, Criado et al., 2007b), this finding implies that the N-A-S-H gel formed in the AAFA with microwave curing could be more Al-rich than that formed with conventional thermal curing.

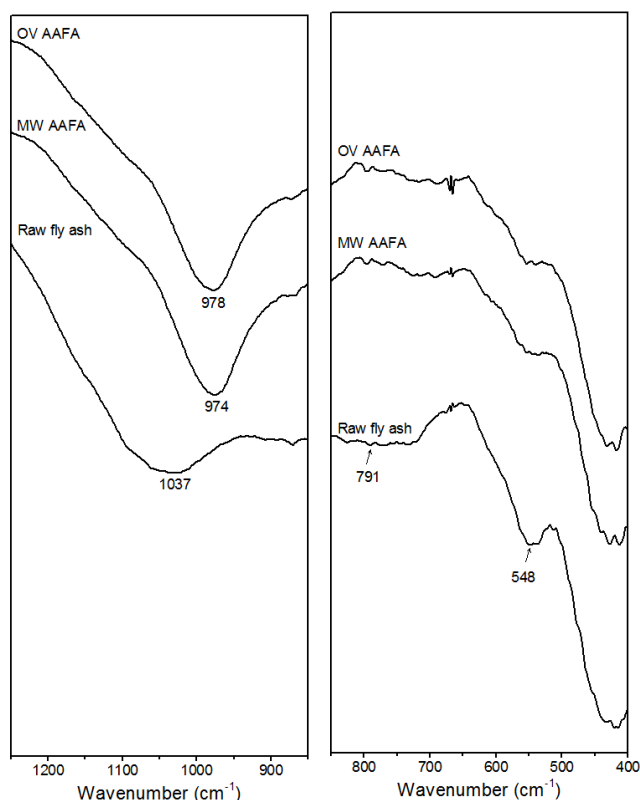


Fig. 4. FTIR spectra of raw fly ash and AAFA samples

^{29}Si NMR

The ^{29}Si NMR spectra of the raw fly ash and the two AAFA samples are illustrated in Figure 5. The deconvolution of the spectra can reveal different components in the fly ash and AAFA samples. In the raw fly ash, the peaks detected at -85.0 ppm, -93.3 ppm, -98.3 ppm, -101.6 ppm and -105.3 ppm can be attributed to the different Si environments in the amorphous phase (Fernandez-Jimenez et al., 2006b, Criado et al., 2008). After alkali activation, the broad resonance band in the raw fly ash shifted in both AAFA samples, indicating the reaction products formed in the AAFA samples (Palomo et al., 2004, Criado et al., 2008). The chemical shifts which can be ascribed to various $\text{Q}^4(\text{mAl})$ units in AAFA samples are given in Table 3. It can be seen that the chemical shift of $\text{Q}^4(4\text{Al})$ and $\text{Q}^4(3\text{Al})$ in MW AAFA are less negative than that in OV AAFA, indicating again more AlO_4 units were involved in the structure of N-A-S-H gel.

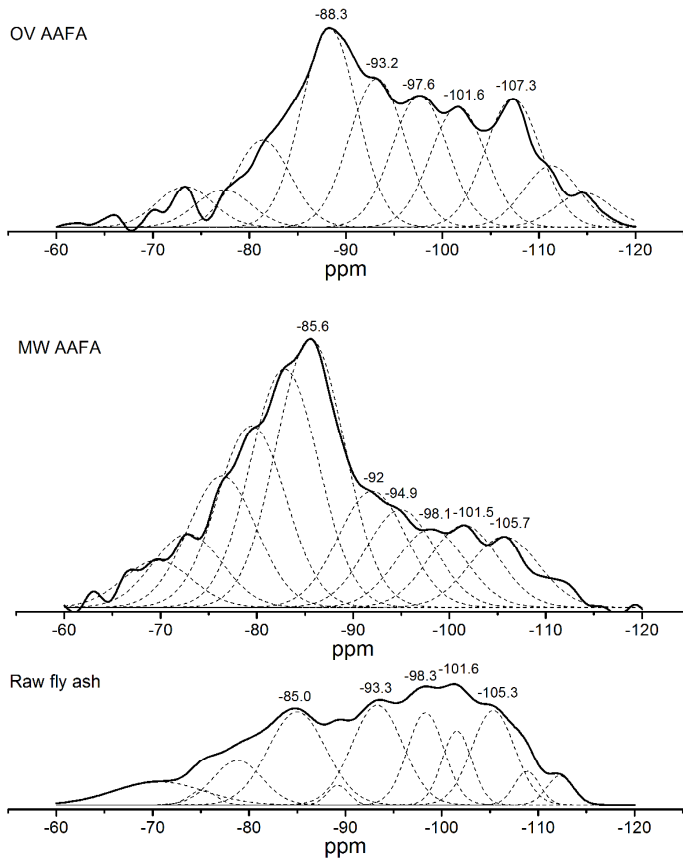


Fig. 5. ^{29}Si NMR spectra of raw fly ash and AAFA samples

Table 3. ^{29}Si signals of N-A-S-H gel formed in OV AAFA and MW AAFA

Sample	$\text{Q}^4(4\text{Al})$	$\text{Q}^4(3\text{Al})$	$\text{Q}^4(2\text{Al})$	$\text{Q}^4(1\text{Al})$	$\text{Q}^4(0\text{Al})$
		-88	-93/-94	-98	-103/-104
OV AAFA	-88.3	-93.2	-97.6	-101.6	-107.3
MW AAFA	-85.6	-92/-94.9	-98.1	-101.5/-105.7	-

MIP

Figure 6a shows the curves of cumulative volume of mercury intruded into the two AAFA samples, which is in line with the results given in Table 4 that OV AAFA obtained a higher total porosity (32.11%) than MW AAFA (20.99%). Figure 6b gives further details of the pore size distribution of the two AAFA samples. It can be seen that there are two peaks representing the ranges of the pores in the OV AAFA, which could be assigned to gel pores (10-140 nm) and capillary pores (140-1000 nm) respectively (Ma et al., 2013). However, most of the pores in the MW AAFA only fell in one range (140-1000 nm), representing the capillary pores. As indicated in Figure 6b, with similar capillary pore distribution between OV AAFA and MW AAFA, the main reason contributed to the higher porosity of OV AAFA is the large amount of gel pores in OV AAFA.

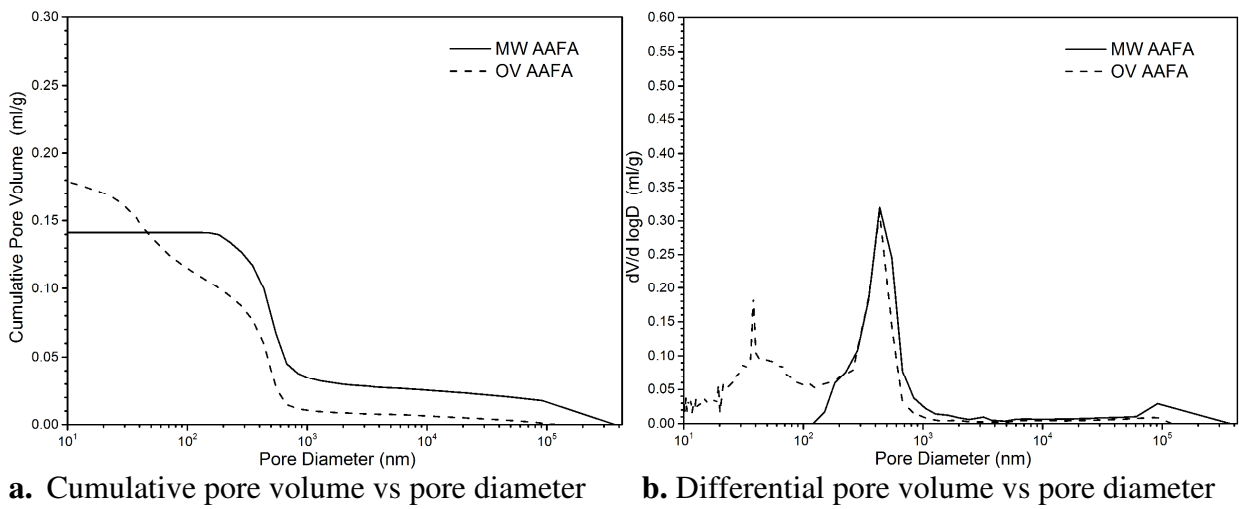


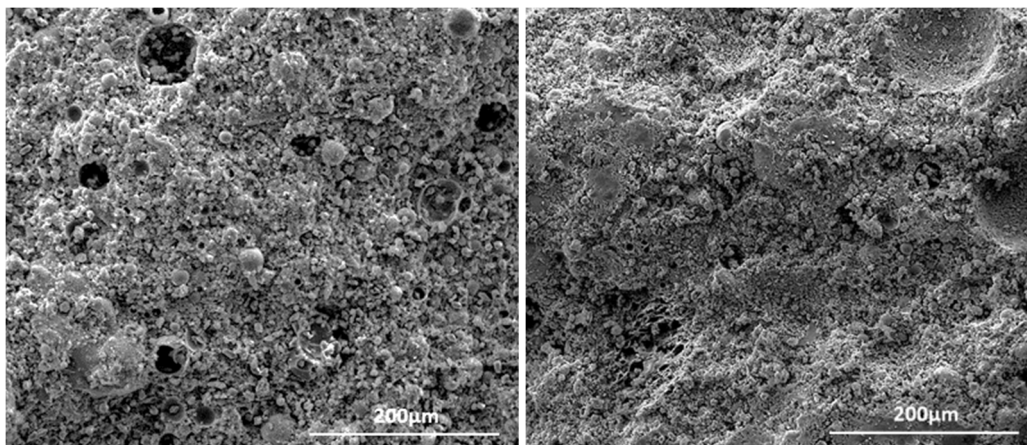
Fig. 6. Pore size distributions of AAFA samples

Table 4. Total porosities of OV AAFA and MW AAFA

Sample	Total porosity
OV AAFA	32.11%
MW AAFA	20.99%

SEM

Figure 7 presents the fractured surface of the two AAFA samples manufactured with different curing methods. Both of them featured heterogeneous microstructure. The OV AAFA showed a more porous structure in comparison with MW AAFA. The reaction products formed on the surface of fly ash particles and in the region between the fly ash particles were also observed in both samples.



a. OV AAFA **b. MW AAFA**

Fig. 7. SEM images of AAFA samples

DISCUSSIONS

In general, alkali activation of fly ash is considered to consist of two stages, namely, dissolution and gelation (Fernandez-Jimenez et al., 2005). It has been agreed that the external heat is essential for the activation of fly ash, particularly with NaOH as the activator. This could be explained by the activation barrier which needs to be overcome with the assistance of external heat before the activation can take place (Bakharev, 2005b). Conventional thermal heating is a slow process relying on heat conduction, whilst microwave can heat up dielectric materials rapidly and volumetrically. The dissolution process can thus be sped up and the reaction rate could be increased by microwave curing due to the high heating efficiency, which has been verified by the higher compressive strength obtained by MW AAFA sample. On the other hand, it seems microwave can also promote the gelation of AAFA as the MW AAFA formed a more compacted microstructure than OV AAFA, which is in line with the strength results.

N-A-S-H gel, the primary reaction product, contributes to the strength development of AAFA. The strength developed is in proportion to the N-A-S-H gel formed. In the previous research, it was found that the N-A-S-H gel formed in the early stage of alkali activation is Al-rich (Gel 1) and a more stable Si-rich N-A-S-H gel (Gel 2) is formed with more silica groups taking part in the gelation process. Compared to Gel 1, the increase of Gel 2 formed is closely related to the mechanical strength rise (Fernandez-Jimenez et al., 2006b). However, both FTIR and NMR results have shown that the N-A-S-H gel formed in MW AAFA is more Al-rich than that formed in OV AAFA, which implies that AlO_4 units may have the advantage of absorbing microwave energy over SiO_4 units, leading to the formation of Al-rich gel. Nonetheless, it seems that the microwave can facilitate the dissolution of fly ash particles and the formation of denser microstructure which could have compensated the weak bond of Al-rich gel. However, the possible selective absorption of AlO_4 units and SiO_4 units to microwave needs further investigation.

CONCLUSIONS

The following conclusions can be drawn from this study:

- With similar temperature history, AAFA manufactured with microwave curing exhibited higher compressive strength, but less energy consumption than that with the conventional thermal curing.
- The microwave curing favoured the formation of Al-rich N-A-S-H gel in AAFA and a more compacted microstructure.

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