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# Mixed hydrogen and biofuels cement clinker: Characterisation, microstructure, and performance

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# ABSTRACT

Over 35 % of the CO<sub>2</sub> associated with cement production comes from operational energy. The cement industry needs alternative fuels to meet its net zero emissions target. This study investigated the influence of hydrogen mixed with biofuels, herein designated net zero fuel as an alternative to coal, on the clinker quality and performance of cement produced in an industrial cement plant. Scanning electron microscopy, X-ray diffraction and nuclear magnetic resonance were coupled to study the clinker mineralogy and polymorphs. Hydration and microstructure development in plain and slag blended cements based on the clinker were compared to commercial cement equivalent. The results revealed a lower alite/belite ratio, but a significant proportion of the belite was of the  $\alpha'_{\rm H}$ -C<sub>2</sub>S polymorph. These reacted faster and compensated for the alite/belite ratio. Gel and micro-capillary pores were densified, which reduced total porosity and attained comparable strength to the reference plain and blended cement. This study demonstrates that the investigated net zero fuel-produced clinker meets compositional and strength requirements for plain and blended cement, providing a feasible pathway for the cement industry to lower its operational carbon significantly.

#### 1. Introduction

Concrete is the most used manufactured material. The production of cement, the binder that holds the aggregate and affords concrete its desirable properties, is energy and CO<sub>2</sub>-intensive. Approximately 3.2 GJ energy is required to produce one tonne of cement clinker in a modern kiln [1], and the process generates 0.58–0.87 tCO<sub>2</sub>e/t of cement [2]. This accounts for over 35 % of the operational carbon from cement kilns, with the remaining 65 % associated with the breakdown of limestone into quicklime [3], which is inherent in cement clinker manufacturing. For the cement industry to attain its 2050 emissions reduction targets, coupling approaches to lower energy-related emissions and increasing the use of alternative materials is essential.

Cement decarbonisation strategies have focused on reducing the clinker factor through increased use of supplementary cementitious materials (SCMs) [4], alternative fuels [5] and carbon capture utilisation and storage (CCUS) from the cement plant [6]. CCUS and carbon-free clinkering raw materials will be game changers, but the enabling

technologies are still evolving and remain medium to long-term strategies. Meanwhile, blended cements, which reduce the clinker factor (and offset the related emissions) in concrete, have been standardized and deployed for many decades. However, slow strength evolution limits practical clinker substitution to at most 50 % in most applications [4,7]. Moreover, the closure of coal combustion power stations and the shift from blast furnaces to electric arc furnace steel production threaten the availability of suitable industrial SCMs, e.g., pulverized fuel ashes and ground granulated blast furnace slag (GGBS), putting into doubt the sustainability of emission reduction through conventional SCMs. Consequently, carbon-free fuels for clinker production will be a practical and impactful strategy towards net zero emission goals.

The energy requirements in the cement plant are traditionally met by coal, petcoke or natural gas, with air as the oxidiser. Increasingly, liquid and solid wastes with different calorific values are used in industrial cement plants as cost and environmental management strategies. Alternative fuels in the cement plant reduce the demand for fossil fuels and energy-related emissions. For example, several cement plants have

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utilised municipal and industrial wastes, sewage sludge, scrap tyres and biomass residues, providing 20–50 % of the cement plants' operational energy [8,9]. However, these fuels differ in carbon, moisture, trace elements and volatile contents, which affect their burnability [10], modifying the kiln's temperature profile and, consequently, the clinker composition and emissions from the plant [11].

The modification of the clinker composition and the availability and cost of fuel are significant factors influencing fuel switching in the cement plant. Commonly used alternative fuels, while having lower calorific values than fossil fuels [5,12], present unique challenges. Their use is associated with a lower heating rate and, depending on the fuel mix, can reduce the peak temperature in the kiln [5]. The effect of such changes on the clinker quality can be profound. For example, moderate reductions in the heating rate were associated with under-burning, where the clinkers were characterised by a low alite/belite ratio and more free lime [13]. Conversely, a higher heating rate increased the  $M_1/M_3$  alite polymorphs ratio with larger alite crystals due to the dissolution of lime and belite [14]. Volatiles in the fuels also affect the clinker polymorphs. For example, phosphorus in agriculture waste and meat and bone meal (MBM) can also be incorporated into the calcium silicate minerals. This stabilises belite over alite, as reported in Refs. [15–17] and elevates the free lime content in the clinker [18]. Similarly, metals in scrapped tyres used as alternative clinkering fuel increase zinc concentration in the resulting clinkers [19]. Meanwhile, alterations in alite grain sizes and polymorphs were identified in biomass and tyre-derived fuel clinkers, which reduced reactivity [20]. Besides trace element inclusions, relationships between the cement kiln's temperature profile and the reactivity of the calcium silicate polymorphs have been reported. The alite polymorphs were shown to increase reactivity in the order  $M_3 < M_1$  [21] and for belite,  $\gamma < \alpha'_L < \beta < \alpha'_H < \alpha$  [22,23]. Therefore, it is imperative to identify potential polymorphic changes in the main clinker phases arising from changes in the kiln fuel composition and, particularly, the switch to new cleaner fuels.

Hydrogen-derived fuels have attracted attention across several industries due to their high calorific value, combustibility and zero carbon intensity. Hydrogen may be generated from natural gas with or without carbon capture, classified as blue and grey hydrogen, respectively [24] or by electrolysis using renewable energy [25]. Presently, supplies to the hydrogen-intensive petroleum and agrochemical industries are met through grey sources. The feasibility of hydrogen-powered cement kilns, integration and synergies for hydrogen generation [26] and their combination with other fuels [10] have been modelled in the literature. For example, the integration of a Cu-Cl thermochemical cycle in cement works as a low-cost and low-energy alternative to water electrolysis via electrification was analysed in Ref. [27]. The regenerative loop exposed cupric chloride salts to steam to form copper oxychloride, which melted at 500 °C, generating hydrogen and oxygen and recycling the chloride salt. Hydrogen from the decomposition of liquefied ammonia was analysed in Ref. [28]. A portion of the thermal energy from the kiln's reaction was used to decompose ammonia, the conversion rate of which exceeded 80 % at 450 °C depending on the catalyst. At the same time, the bulk composition of the resulting clinker was comparable to the coal-produced clinker except for slightly elevated alkali levels. Despite the promising hydrogen production pathways, an understanding of the impact of faster combustibility on clinker quality is lacking. A recent simulation study [10] identified potential fuel miscibility and temperature distribution issues associated with alternative cement kiln fuels including blends with hydrogen. Consequently, industrial-scale trials of such fuels and systematic assessment of the clinker produced from them are imperative to optimise fuel composition and performance.

In this study, a systematic experimental programme is followed to investigate representative clinker and cement from a recently completed net zero fuel switching (i.e., mixed hydrogen and biofuels) trial in the UK. The objective is to examine the influence of the fuel switching on the clinker minerals. Further, hydration of plain and blended slag cements based on the net zero fuel clinker is studied alongside the microstructure and strength development to elucidate underlying factors controlling performance of the cements compared to fossil fuel equivalent.

# 2. Materials and methods

# 2.1. Materials

A net zero fuel produced cement clinker and its corresponding CEM I 52.5 N from a recent UK fuel switching trial project [29] was characterised. The fuel comprised up to 20 % hydrogen mixed with 35 % glycerine and 10 % MBM partially replacing coal in the cement pre-calciner and kiln. Hanson Cement UK supplied the clinker and cement.

Four cements, namely the net zero fuel CEM I 52.5 N ( $C_{NF}$ ), a reference commercial grade CEM I 52.5 N ( $C_{FF}$ ) and their corresponding blends with GGBS designated  $C_{NF}$ S and  $C_{FF}$ S, respectively were investigated to assess the impact of the net zero clinkering fuel on hydration, microstructure, and strength development.

In preparing the blended cement, a 1:1 ratio was maintained between the CEM I and GGBS and the total sulphate content was adjusted to 3 %. Particle size distributions of  $C_{\rm NF}$ ,  $C_{\rm FF}$  and GGBS (S) were measured using a Malvern Panalytical Mastersizer 3000 with an Aero S air dispersion unit and standard Venturi dispenser. Values of refractive index 1.680 and absorption coefficient 0.1 were used to process the data. Three portions of each sample were analysed, and the results were averaged to produce a volume and cumulative distributions as shown in Fig. 1. The Blain fineness was 492, 475 and 510 kg/m<sup>2</sup> for  $C_{\rm NF}$ ,  $C_{\rm FF}$  and GGBS respectively.

For determining the oxide compositions, the as-received cement and GGBS were initially tested for loss on ignition (LOI) at 900 °C in a muffle furnace before the powders were prepared for X-ray fluorescence (XRF) using the fused bead method. The results are shown in Table 1.

Compositionally, the two cements were similar except for slightly elevated levels of alkalis in  $C_{NF}$  and higher LOI reflecting the higher limestone content than  $C_{FF}$  cement. Meanwhile, the  $P_2O_5$  content in  $C_{NF}$  was lower than expected for clinkers made with MBM-containing fuels. However, at 10 % fuel substitution with MBM, the detected 0.2 %  $P_2O_5$  was acceptable. It should also be noted that phosphorous incorporation into clinkers, a risk associated with high MBM as cement kiln fuel substitute, depends on its concentration and the feed rate [30].



Fig. 1. Particle size distribution of  $C_{NF}$ ,  $C_{FF}$  and GGBS as determined by laser granulometry.

#### Table 1

Chemical composition of CNFF, CFF and GGBS as measured by XRF (% weight).

Material	CaO	SiO <sub>2</sub>	$Al_2O_3$	Fe <sub>2</sub> O <sub>3</sub>	MgO	K <sub>2</sub> O	Na <sub>2</sub> O	$SO_3$	TiO <sub>2</sub>	$P_2O_5$	C1	ZnO <sub>2</sub>	LOI
C <sub>NF</sub> C <sub>FF</sub>	63 64.9	19.5 19.5	4.9 5.1	2.1 3.1	2.3 1.1	0.6 0.5	0.3 0.13	3.5 3.1	0.2 0.4	0.2 0.1	0.07 0.05	0.03 0.01	3.3 1.9
GGBS(S)	41.8	34.9	11.6	0.5	5.8	0.5	0.1	3.1	1.1	0.02	-	-	1.5

#### 2.2. Methods

The study comprised two experimental stages.

- 1. Characterisation of the anhydrous net zero fuel (NF) clinker and cement.
- Characterisation of reaction kinetics, microstructure and performance of plain and GGBS blended cement based on the net zero fuel clinker in comparison to the reference commercial cement equivalent.

2.2.1. Characterisation of the anhydrous net zero fuel clinker and cement The as-received clinker was prepared for microanalysis under a scanning electron microscope (SEM), while the anhydrous cement and corresponding enriched silicate and interstitial phases were investigated using X-ray diffraction (XRD) and solid-state nuclear magnetic resonance (NMR). The procedures for selectively enriching the silicate and interstitial phases in the cements were adopted from Ref. [31]. For enriching the silicates, the cement was dissolved in potassium hydroxide and sucrose extraction solution prepared at 9:1:1 in 300 ml deionized water at 95 °C. After 1 min of stirring, the solution was vacuum filtered using an 11 µm cellulose filter paper in a Buchner funnel. The residue was washed with deionized water and methanol before drying at 60 °C. Meanwhile, the interstitial phases were enriched by dissolving 5 g of the cement powder in a 1:15 salicylic acid and methanol solution before vacuum filtering and drying the residue at 90 °C. The dry powders were stored in a desiccator until characterisation.

For observation under the SEM, the clinker granules were embedded in resin under vacuum and cured overnight at room temperature. After removal from the mould, the resin was polished down to 0.25  $\mu$ m using silicon carbide paper and cloths with diamond paste. Images were acquired in backscattered electron mode using a Zeiss EVO 15 SEM with Lab6 filament equipped with a 150 mm<sup>2</sup> Oxford Instruments X-Max EDS detector controlled with the Aztec software. The instrument was operated at 15 KeV accelerating voltage. Elemental mapping and point analyses were carried out to investigate the composition of the main clinker phases, with the backscattered images and combined maps used to determine the grain sizes.

XRD data were acquired on the as-received and selectively enriched  $C_{\text{FF}}$  and  $C_{\text{NF}}$  powder samples. The objective was to identify potential differences in the polymorphs of the main clinker phases resulting from the net zero fuel. The measurements were performed on a Phillips Empyrean diffractometer using a CuKα anode operated at 45 kV and 40 mA and equipped with a PIXcel3D detector without a monochromator. The powders were backloaded into a 10 mm diameter sample holder and scanned on the PW3064 spinner stage. Data acquisition was done at 0.0131 step size with 73.6 s per step between 5 and 80  $2\theta^{\circ}$  range. Rietveld analyses of the diffraction data were conducted using the TOPAS Academic v7.21 software. A control file comprising crystallographic information files for the identified minerals, constrained, and refined on the enriched samples' data was used to quantify the crystalline phases in the anhydrous cements. The external standard method was used to determine the absolute phase contents using a 98 % purity corundum standard.

Solid-state magic angle spinning (MAS) NMR measurements were performed to understand the impact of the fuel switch on the chemical environment of Al and Si. Data was acquired on a Bruker Avance III HD 400 MHz wide bore magnet (magnetic field 9.4 T) with all data acquired at 54.74° magic angle. <sup>29</sup>Si MAS NMR spectra were collected at an operating frequency of 79.48 MHz, using a zirconia rotor, and spun at 6 kHz in a 7 mm solid-state MAS probe using a 90° pulse of duration of 5.5  $\mu$ s, a relaxation delay of 30 s and a minimum of 2048 scans. <sup>27</sup>Al MAS NMR spectra were collected at an operating frequency of 104.23 MHz, using a zirconia rotor, and spun at 20 kHz in a 2.5 mm solid-state MAS probe, and employed a 90° pulse of duration 0.23  $\mu$ s, a relaxation delay of 0.5 s and a minimum of 16,384 scans. Chemical shifts in the <sup>29</sup>Si and <sup>27</sup>Al spectra are reported in ppm with reference to external samples of tetramethylsilane. Baseline corrections on the data were done in the Bruker TopSpin 4.2 software, and the <sup>29</sup>Si spectra were simulated and deconvoluted using the Voight line profile fitting in the Origin Pro software to quantify the alite and belite fractions.

# 2.2.2. Hydration, microstructure, and performance of plain and blended cements as a function of clinkering fuel

The cement compositions used for the hydration, microstructure and performance studies are shown in Table 2. The anhydrous blends were homogenised in a roller ball mill using polymer balls to avoid further grinding. All samples were investigated at 0.5 w/c ratio. For characterising the microstructures, paste samples were prepared according to the procedure for mortars in EN 196-1 [32], but without aggregate. These were investigated for hydration kinetics, phase assemblages and pore structures using isothermal calorimetry, SEM, XRD, thermogravimetric analysis (TGA), mercury intrusion porosimetry (MIP) and dynamic vapour sorption (DVS).

Isothermal calorimetry was used to assess the impact of the net zero fuel on hydration kinetics in plain and GGBS blended cements. Measurements were performed on 9 g of paste prepared at 0.5 w/b ratio using deionized water. The heat signal was corrected with quartz reference samples at the same water to solid ratio. The heat of reaction was monitored for 7 days in a TAM Air 8-channel calorimeter and the data presented in terms of the reaction rate and cumulative heat normalised by the mass of anhydrous cement.

Samples for XRD, SEM, TGA, MIP and DVS were cast into 15 ml plastic vials, sealed, and rotated on a low-speed planetary rotator for 12 h to prevent bleeding. Still under sealed conditions, the samples were stored in a water bath maintained at 20 °C until hydration stoppage. After 1 and 28 days of hydration, specimens for XRD and TGA were crushed to pass 63  $\mu$ m sieve, and hydration stopped using the double solvent removal technique [33]. Following the hydration stoppage, the samples were stored in mini-grip bags in a glovebox under a nitrogen atmosphere until they were tested.

The XRD data acquisition protocol described above for the anhydrous cement was used to scan the hydration-stopped samples after 1 and 28 days. The weight fractions of residual clinker phases were

Table 2	
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Composition of mixes investigated	(%).
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Mix Id	Constituent								
	CEM I 52.5 N (Ref)	CEM I 52.5 N (Net-zero mixed fuel)	GGBS (S)	Anhydrite					
C <sub>NF</sub>		100	-	-					
C <sub>FF</sub>	100	-	-	-					
C <sub>NF</sub> S	-	51	47	2					
C <sub>FF</sub> S	51	_	47	2					

quantified and compared to the anhydrous for determining the degree of clinker hydration. Meanwhile, the Rietveld/PONKCS method, described in Ref. [33], was used to quantify the residual GGBS content and hence the degree of hydration of GGBS in  $C_{\rm NF}S$  and  $C_{\rm FF}S$ . The control file was expanded to include crystallographic files for portlandite (ICSD #43433) [34], hemicarboaluminate (ICSD #263124) [35] mono-carboaluminate (ICSD #59327) [36], hydrotalcite (ICSD #133742) [37], ettringite (ICSD #155395) [38] and a model for C-S-H based on the tobermorite structure of Richardson [39] for quantitative analyses of the hydrated phase assemblages, which were normalised per 100 g of the anhydrous binder.

The 28 days hydrated cement samples for SEM were 2 mm thick slices cut from the paste cylinders using a low-speed Isomet cutter. These were hydration stopped by soaking in isopropanol for 48 h and rinsed in diethyl ether, resin impregnated and polished. Backscattered electron images, elemental mapping and point analyses were carried out to investigate the composition of the main reaction product, the calcium silicate hydrate (C-S-H) using the same SEM and data collection protocols described above for the clinker granules.

TGA data were acquired on a Netzsch thermal analyser 449 F5 Jupiter using platinum sample holders. For each measurement, about 25 mg of ground powder sample was heated at 20 °C/min up to 1000 °C in a nitrogen-purged atmosphere at a 60 ml/min gas flow rate. The mass change and its derivative were used to identify the water-rich hydrated phase assemblages. The bound water and portlandite contents were calculated between 50 and 550 °C and ~400–500 °C from the TGA curve using equations (1) and (2), respectively. The calculated weight fractions were normalised to the ignited weight at 1000 °C.

$$W_n(\%) = \frac{(M_{50^\circ C} - M_{550^\circ C})}{M_{550^\circ C}} * 100\%$$
(1)

$$CH(\%) = \frac{CH_{TG}^{74/18}}{M_{550^{\circ}C}} * 100\%$$
<sup>(2)</sup>

Where,  $W_n$  is the bound water in the sample, *CH* is the portlandite content; *CH*<sub>TG</sub> is % weight loss from water associated with portlandite, calculated from the tangent method,  $M_{50^\circ C}$  is the ignited weight at 50 °C;  $M_{550^\circ C}$  is the ignited weight at 550 °C.

MIP measurements were performed on 1–2 mm crushed, and hydration stopped cement samples using a Quantachrome Instruments' PoreMaster-60. The data acquisition protocol and analysis are detailed elsewhere [40]. The MIP data are presented in terms of the cumulative intruded volume and the derivative.

A separate fraction of the crushed samples was used for DVS measurement on a nitrogen-purged Surface Measurement System DVS Resolution. The relative humidity (partial pressures) from 0 to 95–0 in 5 % steps at 25 °C with the time per step set to achieve equilibration at each step, using a dm/dt value of 0.002 %/minute and a minimum step time and stability period of 10 min. Data was analysed using the SMS DVS Standard Analysis suite of Microsoft Excel macros to calculate an isotherm from the raw data. The Isotherm Analysis suite was used to calculate the BET surface area. Other models used are the Dubinin-Radushkevich (DR), micropore analysis (t-plot) [41] and a modified Barrett, Joyner, and Halenda (BJH) model for mesopore analysis [42]. The micro and mesopore analysis calculations were based on a comparison with a non-porous reference material; this was obtained by running a sample of zirconium silicate under the same conditions as the hydrated cement samples. The DVS data is presented in terms of the micro and meso pore distribution after normalising the isotherms by the sample mass at 5 % RH.

Mortar samples for compressive and flexural strength testing were  $40x40 \times 160$  mm prisms, prepared at 1:3:0.5 binder to sand and water ratios according to EN196-1 [32] and tested after 1, 7 and 28 days. At each age, three prisms were tested for flexural strength and the resulting six halves were tested for compressive strength. The specimens for the

1-day strength were tested immediately after removal from the mould, while those for 7 and 28 days were cured in a water bath maintained at 20  $^\circ\text{C}.$ 

The phase assemblages formed as a function of time and degree of hydration of the cement clinker minerals and GGBS were verified by thermodynamic modelling using the GEMS Selektor v3 package. The thermodynamic data were taken from the PSI– Nagra [43] and Cemdata 18 [44] databases. The mineral composition of the cement as determined by QXRD and bulk composition of GGBS from XRF were used as inputs. Dissolution of the clinker minerals and GGBS as determined by QXRD/PONKCS were fitted using the empirical phase reaction model of Parrot and Killoh [45] with the coefficients adjusted as suggested in Ref. [46].

# 3. Results and discussion

# 3.1. The net zero fuel clinker and corresponding anhydrous CEM I

Fig. 2a shows backscattered electron micrograph of the unground clinker and the overlayed elemental maps of sulphur, magnesium, iron, silicon, and aluminium is shown in Fig. 2b. Granules of free lime were not observed, which is indicative of complete burning of the raw meal. The major clinker phases can be identified alongside variations in the grain sizes and textures of the calcium silicate phases while arcanite and periclase occupied the dark patches in the micrograph. However, the belite to alite contrast was less distinct than those reported in Ref. [47] where polished specimens were etched for observations under light microscope. Nevertheless, spherical and larger dendritic belite grains were evident from the overlayed elemental maps and were characterised by fine textured nodules, while the alite crystal's characteristic micro-cracks were noticed. Similarly textured belite, distinct from the lamellae crystals was reported elsewhere [48] and attributed to recrystallization reactions during polymorphic transitions and quenching of the clinkers [49]. The texturing of belite was also shown to be favoured at higher Al/Fe ratios [22]. Fig. 2b further shows lower iron incorporation into the silicates such that the Al/Fe atomic ratio measured in the belites was  $\sim$  3.6, being greater than the ratios reported in Ref. [22], where Al/Fe ratios were below 2.3, but the observation was comparable to average ratios reported in Ref. [50]. This implies that factors other than the Al/Fe ratio may explain the observed textures. Meanwhile, the interstitials nucleated at the silicate boundaries, as expected with the aluminates and ferrites distinguishable through the combined elemental maps.

The shapes and sizes of the calcium silicate minerals can reveal important information about the kiln temperature profile and potential fuel-raw meal interactions. The measured average grain sizes of alite ranged between 10 and 30  $\mu$ m while belites were 15–40  $\mu$ m, consistent with those reported in the literature [50]. The dendritic features could be due to dissolution of belites, probably constrained by limited lime for conversion into alite. The features found on the micrograph in Fig. 2a and b, therefore, highlight the specific effects of the fuel composition and temperature profile on the clinker minerals, which bears significant implications on their grindability and subsequent reactivity of the cement.

Elemental compositions of the main clinker phases based on atomic ratios from EDS point analyses (shown in Table 3) offer further insight into the modifications which underpinned the polymorphic differences in the clinker. The reduction in alite's calcium concentration was balanced with sodium, potassium, and magnesium, while slightly more aluminium and phosphorous were incorporated into both alite and belite, which also contained lower iron than typically reported [50,51]. These inclusions partly explain the observed texturing in the micrographs in Fig. 2. The interstitials meanwhile showed a slight increase in the aluminium substitution by silicon and iron in the aluminate and ferrite phases. It was not possible to distinguish the aluminate polymorphs due to resolution of the EDS point analysis technique. However,



**Fig. 2.** BSE images of the unground clinkers from the (a) net zero fuel and (b) overlayed elemental maps. Note: In (b), blue = sulphur, purple = magnesium, red = iron, light blue = silicon and green = aluminium. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

### Table 3

Stoichiometric composition of the major phases in the  $C_{\rm NF}$  clinker based on the atomic ratios from EDS analysis.

Phase	Composition
C <sub>3</sub> S	$(Ca_{0.85}Mg_{0.04}K_{0.01}Na_{0.01}Fe_{0.009})_3(Si_{0.9}Al_{0.04}P_{0.01}S_{0.04})O_{4.6}$
C <sub>2</sub> S	$(Ca_{0.96}Mg_{0.016}K_{0.011}Na_{0.012})_2(Si_{0.9}Al_{0.062}Fe_{0.017}P_{0.003}S_{0.02})O_{3.8}$
C <sub>3</sub> A	$(Ca_{2.76}K_{0.03}Na_{0.02}Mg_{0.2}Ti_{0.04})(Al_{0.96}Fe_{0.42}Si_{0.28})O_{5.86}$
C <sub>4</sub> AF	$Ca_{2.03}Al_{0.98}Fe_{0.5}Mg_{0.17}Si_{0.25}Ti_{0.03}O_{4.7}$

the 8 % and 42 % substitution of calcium and aluminium respectively in the  $C_3A$  indicate a mixture of cubic and orthorhombic  $C_3A$ .

The as-received  $C_{NF}$  sample and the corresponding enriched samples were analysed by XRD and NMR to examine the implications of the elemental variations on mineralogy. Fig. 3a shows diffraction patterns of the enriched silicate minerals and the enriched interstitial in Fig. 3b, focused on the relevant angular windows to reveal characteristic peaks and the effect of minor inclusions. X-ray reflections denoting the major clinker phases can be identified, albeit with some structural modifications evident from the peak shapes, particularly the intensity ratios of the primary *hkl* reflections. The doublet reflections between 32 and 33° 20 in Fig. 3a indicate the monoclinic form of alite but, the overlap with belite in the (60 $\overline{6}$ ) and (224) reflection ranges causes uncertainties in the polymorph assignment. The relative intensity ratio between the (60 $\overline{6}$ ) and (224) reflections was much greater while the ratio between the (14.24) and (040) reflections observed between 51 and 52.5° 20 exceeded unity. These indicate a mixture of the  $M_1$  and  $M_3$  alite polymorphs [50,52], consistent with the presence of periclase in Fig. 3b and the EDS composition in Table 3, which showed higher magnesium incorporation in alite. Structure search and match on the data confirmed dominance of the  $M_3$  alite with small levels of the  $M_1$  polymorphs. The refined lattice parameters of these matched closely the structures of De La Torre et al. [53] and de Noirfontaine et al. [54] respectively. This implies that the alite polymorphs formed in the  $C_{\rm NF}$  clinker conformed to those typical in industrial cements.

Variations in the belite polymorphs in the  $C_{NF}$  clinker were also evident from Fig. 3a. Assigning XRD peaks to belite in industrial cement is not straightforward due to the substantial overlap with alite in the same 20 range. However, high-temperature XRD data [50] and synthetic belite doped with phosphorous [17] or boron [55] help to identify the belite polymorphs. From Fig. 3a, the *hkl* reflections at (611) and (031) represent  $\alpha'_{H}$ -C<sub>2</sub>S while those at (120) and (121) represent  $\beta$ -C<sub>2</sub>S, thus indicative of the co-existence of the  $\alpha'_{H}$  and  $\beta$ -C<sub>2</sub>S polymorphs in the C<sub>NF</sub> clinker. The former plausibly arose from the higher levels of MBM, which, together with hydrogen and glycerine, replaced coal as fuel. From Fig. 3b, the C<sub>3</sub>A polymorph and C<sub>4</sub>AF were unaffected by the net zero fuel; cubic C<sub>3</sub>A (440) with some orthorhombic C<sub>3</sub>A (224) were observed while the *hkl* reflections at (200, 141) and (002) denoted the



Fig. 3. XRD trace of the selectively enriched  $C_{NF}$  cement: (a) silicate phases after dissolving in the KOH/sucrose solution (b) interstitials after dissolving in salicylic acid/methanol solutions using the methods of Guttridge [31].

C<sub>4</sub>AF phase. Notably, the C<sub>3</sub>A/C<sub>4</sub>AF intensity ratio was more significant in the net zero fuel than typically reported in the literature, suggesting a lower ferrite phase consistent with the EDS maps and elemental ratios in Table 3.

The clinker content and ratios between the main phases are important quality indicators of cement performance. From the results in Table 4, the total clinker content in the plain cement was 90 % while the minor additional constituents comprising a mixture of anhydrite, bassanite, calcite and periclase constituted 10 %. Of the clinker content, alite and belite made up 80 % at 3:1 alite/belite ratio, which is slightly lower compared to typical industrial cement. The  $\alpha'_{H}$ -C<sub>2</sub>S polymorph constituted 55 % of the total belite content, while the  $\beta$ -C<sub>2</sub>S polymorph was approximately 45 %. Occurrence of the  $\alpha'_{H}$ -C<sub>2</sub>S is noteworthy given its faster reaction than  $\beta$ -C<sub>2</sub>S [23,55]. Whether the temperature profile or the fuel composition dictated the observed belite polymorphs is uncertain. However, according to the literature,  $\alpha'_{H}$ -C<sub>2</sub>S which forms at higher temperature than  $\beta$ -C<sub>2</sub>S (i.e. >1160 °C) is stabilized at room temperature through partial substitution of silicon by phosphorous [16] which can be expected with the MBM in the fuel. Increasing the phosphorous concentration was directly correlated with the stabilization of belite over alite [18] and  $\alpha'_{H}$ -C<sub>2</sub>S, at higher phosphorous and iron concentrations in the raw meal [17]. It is noteworthy that limited  $\alpha'_{H}$ -C<sub>2</sub>S formed in clinkers doped with <1.1 % P<sub>2</sub>O<sub>5</sub> but excess phosphate also hindered  $\alpha'_{H}$ -C<sub>2</sub>S formation and retarded hydration of belite-rich cement [16]. Optimisation of the phosphorous concentration in the fuel and hence the biofuel content is therefore essential to maintain the acceptable alite and  $\alpha'_{H}$ -C<sub>2</sub>S contents in the clinker.

Cubic C<sub>3</sub>A existed alongside the orthorhombic type, consistent with the 6.5 % alkalis including magnesium in the aluminate structure as revealed by the EDS results in Table 3. The overall levels of both C<sub>3</sub>A and C<sub>4</sub>AF were in line with the levels in commercial cements and not modified significantly by the investigated fuel.

To evaluate the chemical environments of alumina incorporated into the minerals, <sup>27</sup>Al NMR scans were performed on the as-received cement, enriched silicate and interstitial samples and the results shown in Fig. 4. Two central transitions between 50 and 100 ppm and -20 - 20 ppm denoting tetrahedral (Al(IV)) and octahedral (Al(VI)) coordinated aluminium respectively are noticed. The deshielded centreband at 96 ppm and the main singular peak at 79 ppm plus various sub-peaks characterised the tetrahedral coordinated aluminium, indicative of some  $Al^{3+}$  partially substituting  $Si^{4+}$  in belite and alite. The octahedral coordinated aluminium showed a symmetrical centre band at -20 to 20 ppm centred at ~ -0.6 ppm, coinciding with that of the interstitials and amplified in the enriched interstitials sample. This implies that another aluminium site, likely to be X-ray amorphous, co-existed in the clinker with similar coordination in the silicates and interstitials, being more concentrated in the latter. The tetrahedral centre band at ~56 ppm was unique to the enriched interstitial sample and distinct from previously reported C<sub>3</sub>A centre-bands at around 80 ppm. The lower frequency shift in the Al(IV) of the interstitial sample therefore indicates greater electron density around the aluminium nucleus compared to those substituted for the silicate minerals. This shielding is consistent with the extent of ion substitution, especially silicon in the interstitials as revealed by EDS (Table 3).

The impact of the net zero kiln fuel on the chemical environment of aluminium could only be evaluated qualitatively owing to the spectral resolution. The Al(VI)/Al(IV) intensity ratios of the silicates were greater, indicative of lower Al substitution in the silicates as observed



**Fig. 4.** <sup>27</sup>Al NMR spectrum (9.4 T, 20 kHz, 16,384 scans) showing central transition for Al ions in the (i) as received cement, enriched (ii) silicates and (iii) interstitials in the  $C_{\rm NF}$ cement and after selective enrichment of the silicates and interstitial phases.

from the EDS-derived stoichiometric compositions (Table 3). Some studies have shown that increasing sulphate incorporation into the silicate phases also raised the aluminium concentration [56], which can ultimately impede the reaction of the silicate phases [57].

Meanwhile, the <sup>29</sup>Si NMR spectra (Fig. 5) reveal distinctive information about the silicon coordination in the as-received clinker, which closely matched the enriched silicate spectra. Silicon ions incorporated into the interstitial phases were not detectable at the resolution of the acquired data. Simulation of the enriched silicate spectra allowed a closer examination of the calcium silicates. These were characterised by a well-resolved resonance at -71.4 ppm corresponding to belite, but polymorphs could not be differentiated from the single resonance. The overlapping resonances in the -78 to -66 ppm range are attributable to alite, whose sub-spectra comprised broad resonances at  $\sim$  -70.4, -73.6 and -75.5 ppm, typical of the M3 alite polymorph. The diffused resolution of the alite sub-spectra indicates ion substitution [58], consistent with the elemental composition in Table 3. Moreover, the peak area ratio of belite/alite was 0.36, combined with the bulk SiO<sub>2</sub> from XRF, and the elemental composition (Table 3) yields 55.4 % and 16.2 % alite and belite, respectively. These are within the typical ranges reported for OPC [50,58,59] and show excellent agreement with the quantitative XRD alite and belite contents in Table 4.

The results presented above on the clinkers and the cements have shown polymorphic differences in the alite and belite phases and their relative contents. The EDS analyses also confirmed more magnesium and phosphorous but less sulphur, aluminium, and iron substitution in the  $C_{\rm NF}$  clinker. Moderate variations in the raw meal and volatiles in the kiln (i.e., pre-existing or emanating from the fuels), fuel ashes or the lack of these, plus the kiln temperature profile, might have contributed to the polymorphic changes in the main phases. For example, hydrogen being

Table 4 Mineralogical composition of the C<sub>NF</sub> cement based on quantitative XRD.

Phase	Mixes									
	C <sub>3</sub> S	$\alpha'_{H}-C_2S$	$\beta$ -C <sub>2</sub> S	$C_3A_{cub}$	C <sub>3</sub> A <sub>orth</sub>	C <sub>4</sub> AF	Anhydrite	Basanite	Calcite	Periclase
C <sub>NF</sub>	56.4	9.8	8.3	4.3	2.9	7.1	3.6	0.6	4.8	1.4



**Fig. 5.** <sup>29</sup>Si NMR (9.4 T, 6 kHz, 2048 scans) spectra of the as received clinker (i) enriched calcium silicates (ii) and the interstitial (iii) phases alongside the optimized simulation (trace on the silicates) and deconvoluted belite (iv) and alite (v) phases  $C_{\rm NF}$  cement and after selective enrichment of the silicates and interstitial phases.

more combustible than coal and MBM can cause a sharper heating gradient [10], exposing the raw meal faster to the high-temperature range due to the shorter ignition distance than that typically encountered in coal-rich kiln fuels. The endothermic reactions involved in the formation of belite, and the interstitials plausibly extended the belite formation temperature range further, consuming quicklime, and modifying the overall belite concentration. Therefore, the limiting step in the fraction of alite quantified in Table 4, which is lower than the levels in some commercial cements, may be due to the depletion of quicklime in the stabilized  $\alpha'_{\rm H}$ -C<sub>2</sub>S rather than the attained temperature range for the fuel mix. Moreover, alternative fuels' combustion can also reduce the oxygen and sulphur concentrations in the kiln [10] and, their incorporation in the clinker phases, which potentially contributed to the observed polymorphs and the alite/belite ratio in the C<sub>NF</sub> clinker.

#### 3.2. Hydration kinetics and microstructure

The preceding sections revealed a moderately lower alite/belite ratio and significantly higher  $\alpha'_{H}$ -C<sub>2</sub>S in the C<sub>NF</sub> clinker and cement. Evaluating the hydration kinetics and microstructures formed in cements based on this clinker is imperative to understanding their performance.

Fig. 6a shows the isothermal heat of reaction measured up to 7 days in the plain and blended GGBS cements from the net zero fuel clinker ( $C_{NF}$  and  $C_{NF}S$ ) compared to the reference cements ( $C_{FF}$  and  $C_{FF}S$ ). Moderate differences in the kinetics of the silicate and aluminate reactions as a function of the clinkering fuel were evident. A shorter induction time and an intense main reaction peak characterised  $C_{NF}$  and  $C_{NF}S$  compared to the  $C_{FF}$  equivalent. However, slopes of the acceleration stage reactions leading to the main silicate peak were comparable between the plain cements,  $C_{NF}$  and  $C_{FF}$ , with the peak reached after about 9 h. The aluminate reaction/sulphate depletion peak, observed as a shoulder on the silicate peak was less noticeable in the  $C_{NF}$  sample, seemingly merged with the silicate peak. This occurred after ~15 h in the reference sample, suggesting potential differences in the sulphate/ alumina consumption in the plain cements.

The blended cements,  $C_{NF}S$  and  $C_{FF}S$  showed a lower rate of reaction due to the 50 % reduction in the clinker factor, although with similar features as the plain cements. The silicate reaction was enhanced in  $C_{NF}S$ , but the time to the main reaction peaks was not modified significantly by the clinkering fuel type nor the presence of GGBS. However, the aluminate peaks showed some differences despite comparable  $SO_3$ contents in the cements (see Table 1). The first aluminate peak occurred 3 h faster in  $C_{NF}S$  compared to  $C_{FF}S$ , but the second aluminate peaks were delayed in both blended cements until ~34 h. The higher surface area and fineness of  $C_{NF}$  may explain the greater silicate and aluminate reactions in the corresponding plain and blended cement, while the higher sulphate content in the blended cements (see Fig. 7) explains the delayed second aluminate peaks in  $C_{NF}S$  and  $C_{FF}S$ .

The kinetic differences notwithstanding, the cumulative heat (Fig. 6b) shows marginally high total heat in the  $C_{NF}$  than the reference  $C_{FF}$  and the  $C_{NF}$ S than  $C_{FF}$ S. Moderately elevated alkali content, plus lower iron [17] and aluminium [57] incorporated into the  $C_{NF}$  clinker, possibly accelerated and enhanced the silicates hydration. Moreover, a faster reaction of the  $\alpha'_{H}$ - $C_2S$  could also contribute to the marginal increase in cumulative heat in  $C_{NF}$  and  $C_{NF}S$  than the reference equivalents.

QXRD/PONKCS analysis was used to follow the hydration reactions and products with a view to elucidating the contributory factors



Fig. 6. Early age hydration of plain and blended GGBS cements from the C<sub>NF</sub> and C<sub>FF</sub> cements (a) rate of reaction (b) cumulative heat as measured by isothermal calorimetry.



**Fig. 7.** Predicted phase assemblages as a function of the dissolution kinetics of (a) the clinker minerals in the  $C_{NF}$  cement and (b) with GGBS in the  $C_{NF}$ S cement over the course of hydration. The solid lines denote the modelled dissolution kinetics of the anhydrous minerals, GGBS and anhydrite. The dashed lines represent the predicted phase assemblages in in GEMS. The phase contents of the anhydrous and phase assemblages calculated from QXRD/PONCKS are shown as scatter points after 1 and 28 days.

underpinning the above calorimetry data. Comparisons between the QXRD/PONKCS results and the thermodynamic modelling of  $C_{\rm NF}$  and  $C_{\rm NF}$ S are shown in Fig. 7a and b respectively. The residual clinker minerals and GGBS determined from QXRD/PONKCS analyses were fitted reasonably by the modified phase dissolution model of Parrot and Killoh [45,46]. These were implemented in GEMS, providing a basis to predict the stable phase assemblages, assuming equilibrium in the dissolved and precipitated species. In the early hours of hydration (<~3 h), alkalis and sulphates dissolve rapidly alongside a gradual alite consumption. After that, alite dissolved faster. This point corresponded to the onset of the acceleration stage of hydration (Fig. 6a) with increased

precipitation of C-S-H, portlandite and hydrogarnet predicted (Fig. 7a and b). Ettringite precipitated alongside these, but its content plateaued once anhydrite was dissolved fully. The time to complete anhydrite dissolution reflected the sulphate balance in the cements. After 1 day, the degree of hydration of alite was 55–65 % in mixes  $C_{NF}$  and  $C_{NF}S$ , respectively, reaching 90–98 % by 28 days. Meanwhile, 19–22 % degree of hydration of belite was measured after 1 day and increased to 50–60 % after 28 days. While the measured and modelled degree of hydration of alite was expected, that of belite was significantly higher than typically reported [52,60], but the data is consistent with the polymorphs identified in the  $C_{NF}$  clinker.



Fig. 8. Backscattered electron micrographs showing types and distribution of phase assemblages in plain and blended GGBS cements from  $C_{NF}$  compared to their  $C_{FF}$  equivalent, measured by SEM: (a)  $C_{NF}$  (b)  $C_{FF}$  (c)  $C_{NF}$  S and (d)  $C_{FF}$  S after 28 days of hydration.

#### 3.3. Microstructure

The thermodynamic modelling results in Fig. 7 (a and b) predicted the C-S-H, ettringite, portlandite and silicious hydrogarnet as the main hydrated assemblages with some monocarboaluminate and hydrotalcite. The quantities of these depended on the binder composition and degree of hydration. The predicted phase assemblages were verified experimentally through SEM, TGA and XRD.

Fig. 8(a–d) shows backscattered SEM micrographs of the hydrated cement pastes after 28 days. These reveal an intermix of hydration products, unreacted clinker, GGBS, and capillary pores. Light grey assemblages denoting portlandite were uniformly interspersed within the C-S-H phases and more dominant in the plain cement,  $C_{NF}$  and  $C_{FF}$ .

The hydrated plain cement showed fewer grains of unreacted clinker phases in sample  $C_{NF}$  (Fig. 8a) compared to  $C_{FF}$  (Fig. 8b). In the latter, partially reacted grains with contrasts of bright phases, corresponding to an intermix of belite and ferrite, can be noticed. In the GGBS blends (Fig. 8c and d), traces of unreacted clinker, especially the bright ferrite phases and GGBS grains, can be identified alongside relicts of the reacted GGBS. Differences in the capillary porosity of the pastes can also be identified in the micrographs, being more dominant in the blended cements. The reference  $C_{FF}$  and  $C_{FF}S$  cements showed larger capillary pores than the corresponding  $C_{NF}$  and  $C_{NF}S$  samples. These observations are consistent with the anhydrous cement particle size distribution (Fig. 1) and the reaction kinetics (Figs. 6 and 7).

The TGA/DTG spectra in Fig. 9 measured on 28-day samples confirmed C-S-H and ettringite, carboaluminates, hydrotalcite and portlandite as the main hydration products. Calcite in the hydrated C<sub>NF</sub> samples is residual from the limestone added as a minor additional constituent (MAC, see Table 4). The C-S-H and ettringite endotherms were comparable in the plain cements, but slightly more portlandite can be observed in C<sub>NF</sub>. Comparison of the plain and blended cement endotherms suggest more C-S-H and or ettringite in the blended cement. Both products consume calcium and reduce the portlandite content with the lower clinker factor. Meanwhile, endotherms of carboaluminates (marked as Hc/Mc) were noticed in all samples but were dominant in the C<sub>NF</sub> and C<sub>NF</sub>S samples due to the excess limestone from the MAC. The TGA calculation inserted in Fig. 9 indicates higher bound water and portlandite contents in the plain C<sub>NF</sub> cement, which agrees with the kinetics data in Figs. 6 and 7. The portlandite contents in C<sub>NF</sub>S and C<sub>FF</sub>S are consistent with the enhanced clinker hydration in the cements (see Fig. 7). However, the TGA-measured portlandite content was much lower in C<sub>NF</sub>S than C<sub>FF</sub>S. The higher degree of hydration of clinker,



Fig. 9. TGA/DTG plots showing the thermal decomposition of the main waterrich phase assemblages in the  $C_{\rm NF}$  plain and blended cements compared to  $C_{\rm FF}$ equivalents measured after 28 days of hydration.

would be expected to contribute more portlandite, yet a lower content was measured in  $C_{NF}S$ . This indicates greater portlandite consumption in  $C_{NF}S$ , which agrees with the degree of GGBS reaction in Fig. 7 and the bound water contents (insert in Fig. 9).

In Fig. 10a, XRD patterns of the anhydrous and hydrated pastes focused in the 8–35°  $2\theta$  range to reveal the crystalline and semicrystalline phase assemblages along with the clinker minerals are presented. These confirm the C-S-H, portlandite and ettringite as the primary assemblages in the plain and blended cements by 1 day, and after 28 days, carboaluminates and hydrotalcite can be observed. The observations confirm the phase assemblages predicted by the thermodynamic modelling in Fig. 7(a and b). The exception is hydrogarnet. Due to its weak XRD reflection and overlap with the C-S-H, distinct peaks corresponding to the latter could not be assigned. Calcite from limestone in C<sub>NF</sub> stabilized ettringite in the plain and blended cement. Slightly more calcium sulphate in the blended cement retarded calcite dissolution and hence carboaluminate precipitation. In the 28d samples, hemi and monocarboaluminate reflections were identified in CNFS but only monocarboaluminate in C<sub>NF</sub>, plausibly due to the limited aluminium availability after the interstitial aluminate depletion.

Quantitative analyses of the XRD data, showing the unreacted clinker minerals, GGBS and phase assemblages formed, plus the calculated degree of hydration of clinker and GGBS are presented in Fig. 10b. Between 1 and 28 days, more portlandite and C-S-H were calculated in both cements, consistent with the dissolution kinetics (Fig. 7a and b) and the bound water (Fig. 9). Compared to C<sub>NF</sub>, the C<sub>NF</sub>S sample showed more ettringite due to the elevated sulphate but less portlandite and C-S-H due to the lower clinker factor and overall degree of hydration respectively. Comparison between the QXRD results and those predicted in GEMS can be seen in Fig. 7(a and b). Consistency between the datasets is remarkable, except hydrogarnet that was not quantified in the analysis and significantly lower C-S-H in the  $C_{\mbox{\scriptsize NF}}S$  samples. Given that hydrogarnet was not quantified, one would expect more C-S-H calculated. It seems the underestimation was at the expense of the residual GGBS calculation by PONKCS, which may imply possibly greater degree of GGBS hydration than calculated.

These results demonstrate that the main hydrates were not affected by the clinkering fuel, but the volume of formed phase assemblages and their evolution with time were modified due to reactivity of the minerals and the MAC in the cement. Stabilization of Hc/MC over ettringite at 28 days and portlandite consumption to form C-S-H and Hc/Mc are wellknown phenomena in plain and blended cement due to the additionally dissolved alumina from GGBS [61] and their interactions with dissolved anionic species [62].

The composition of the C-S-H phase from the plain and blended cements plotted in terms of the Al/Ca versus Si/Ca based on EDS atomic ratios are shown in Fig. 11, together with tabulated Ca/Si and Al/Ca ratios. Note, the Ca/Si ratio was calculated as the reciprocal of the furthest point cluster on the Si/Ca axis while the slope, drawn from the origin through the lowest points on the Al/Ca axis was taken as the Al/Si ratio according to Ref. [39]. Whereas the C-S-H Ca/Si and Al/Si differed between the plain and blended cements, the effect of the clinkering fuel on these was moderate, and the ranges are consistent with the literature on GGBS blended cements [61]. The CNFS and CFFS cements' lower C-S-H Ca/Si ratio is due to a lower calcium content in the GGBS compared to plain cement and the subsequent calcium consumption to form additional hydration products. Meanwhile, elevated alumina in the pore solutions due to GGBS hydration tends to increase the C-S-H Al/Si as in the alite and belite. Consequently, the clinkering fuel had no measurable effect on the composition of the C-S-H phase in the plain or blended cements.

The C-S-H compositional similarity notwithstanding, differences in the reaction kinetics and volume of hydrates affected the pore distribution and porosity. The mercury intruded porosity and its derivative (Fig. 12a) showed three pore clusters ranging between 5 and 20 nm, 20–100 nm and >100 nm, corresponding to the meso, micro-capillary



**Fig. 10.** XRD data showing (a) diffraction trace of anhydrous and hydrated  $C_{NF}$  and  $C_{NF}$ S cements, focused to reveal the main minerals and crystalline phase assemblages and (b) quantitative XRD analysis of the anhydrous cements and pastes after 1- and 28-days hydration. In (a), the bottom set is the trace for anhydrous cements, middle set is hydrated paste after 1-day and top set is hydrated pastes after 28-days. Note, the offsets in the vertical axis of (a) are for clarity. In (b), the degree of hydration of the clinker minerals was calculated for  $C_{NF}$  and  $C_{NF}$ S and GGBS in  $C_{NF}$ S. There is  $\pm 2$  % margin of error associated with the degree of hydration calculated.



Fig. 11. SEM/EDS of the outer product C-A-S-H in the plain and blended GGBS cements based on  $C_{\rm NF}$  compared to  $C_{\rm FF}$  equivalent after 28 days hydration.

and macro-capillary pores. It must be noted that the MIP technique does not resolve pores finer than 4 nm, which dominate the gel and the C-S-H interlayers [63]. In the MIP accessible pore ranges, the clinkering fuel had a negligible effect on the crossover between the meso and capillary pores. The micro-capillary pores were densified in  $C_{\rm NF}S$  and  $C_{\rm FF}S$ , but macro-pores also existed in the corresponding plain cements. For each cement category, however, the  $C_{\rm FF}$  cement showed greater porosity than the  $C_{\rm NF}$  equivalent, but the critical pore size diameters were not modified considerably. Meanwhile, the DVS data (Fig. 12b) gave further insight into the mesopore ranges that were inaccessible by MIP.

The interlayer pores (insert in Fig. 12b) corresponding to the C-S-H interlayer spaces showed slight coarsening in the blended cement. However, adsorption causes swelling and collapse in these pore ranges during the measurement, and hence, it is uncertain whether the aluminium incorporation in the blended cement influenced the trends since the latter were much denser, below  $\sim 1$  nm. Above this, the C<sub>NF</sub> and

 $C_{\rm NF}S$  samples showed a smaller volume of finer pores than the  $C_{\rm FF}$  equivalent. These observations can be explained by the pore refinement arising from the additionally formed hydration products, as revealed by the degree of hydration and weight fraction of hydrates evident from Figs. 9 and 10 respectively.

# 3.4. Compressive strength

Fig. 13(a and b) show compressive and flexural strength of mortars prepared from the investigated plain cements (C<sub>NF</sub> and C<sub>FF</sub>), and their corresponding GGBS blends (C<sub>NF</sub>S and C<sub>FF</sub>S) measured after 1, 7 and 28 days. Higher compressive strength was measured in the  $C_{\rm NF}$  mortars compared to the CFF equivalent, although the difference reduced with time, falling within the measurement error by 28 days. As expected, compressive strength evolved slowly in the blended GGBS mixes, which were still lower than the corresponding plain cement mixes after 28 days. This is consistent with results from strength tests based on similar GGBS [64] and can be explained by the slow reactivity of GGBS compared to cement clinker (see Fig. 10b). Unlike the plain cement mortars, differences in compressive strength of C<sub>NF</sub>S and C<sub>FF</sub>S mortars were within the measurement error, suggesting a negligible influence of the investigated net zero fuel on the strength of blended cement mortars. except at early age when the latter was up to 5 MPa greater. Meanwhile, flexural strength development in the mortars followed similar trends as the compressive strength, comparable between  $C_{\rm NF}$  and  $C_{\rm FF}$  cement whether plain or blended. The strength profiles are consistent with the kinetics and microstructure studies, which identified slightly improved reactivity and densified pore structures in CNF and CNFS cements compared to their fossil fuel equivalent under the same testing conditions.

#### 4. Conclusions and recommendations

This study investigated representative clinker and cement from a net zero-emission fuel that comprised mixed hydrogen, MBM and glycerine used in an industrial cement plant. The hydration, microstructures, and performance of plain and blended cement from the clinker were compared with commercial grade fossil fuel equivalent. Based on the results and discussion, the following conclusions are drawn.



**Fig. 12.** Pore structure of the investigated plain and blended GGBS cement based on  $C_{NF}$  and  $C_{FF}$  cements. (a) Differential and cumulative intruded volume plots from MIP and (b) distribution of micro (insert in 12b) and meso pore distribution calculated from the desorption isotherm as measured by DVS.



Fig. 13. Strength evolution in the investigated plain and blended GGBS cement based on clinkers from net zero and reference. (a) Compressive and (b) flexural strength monitored up to 28 days.

- a. The four main minerals alite, belite, calcium aluminate and calcium alumino-ferrite phases were identified in the net zero fuel clinker although with polymorphic differences prominent in the belite phase. The alite/belite ratio (~3) was slightly lower compared to typical fossil fuel-derived industrial cement clinkers, but the  $\alpha'_{\rm H}\text{-}C_2S$  polymorph was also found in the net zero fuel clinker which constituted about 55 % of the total belite content.
- b. Tetrahedral aluminium coordination (i.e. Al(IV)) in the net zero fuel interstitials was more shielded and pivoted at  $\sim$ 56 ppm compared to 80 ppm reported for Portland cement C<sub>3</sub>A and 81–96 ppm denoting Al(IV) incorporated into alite and belite. Guest ions including silicon in the interstitial phases led to the shielding. Moreover, the octahedrally coordinated aluminium site, likely to be an X-ray amorphous phase concentrated more in the interstitials.
- c. The net zero fuel cement's alite and belite reacted faster and attained an overall greater degree of reaction than the levels reported for fossil fuel produced cement in the literature. The kinetic effects arose from the  $M_3$  alite and  $\alpha'_H$ -C<sub>2</sub>S polymorphs, which reacted faster and

contributed additional reaction products as reflected in the bound water contents.

d. The nature of reaction products including composition of the C-S-H phase, portlandite, ettringite and carboaluminates was not modified significantly by the clinkering fuel, except for the volume of hydrates in the plain and blended cements. The increased kinetics and volume of reaction products reduced porosity compared to the fossil fuel equivalent cements and led to slightly improved strength at the investigated timescales.

The observed polymorphic differences, reaction kinetics and microstructures indicate that the trial cement/clinker fulfils the market requirements, demonstrating a comparable performance to the reference cement. Implementation of hydrogen-based net zero fuels for cement clinker production at an industrial scale is therefore feasible. However, kinetics of combustion of the different fuel substitutes and possible interactions with the raw meal may require optimisation before deploying at scale.

# CRediT authorship contribution statement

Samuel Adu-Amankwah: Writing – review & editing, Writing – original draft, Software, Project administration, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. Ben Douglas: Writing – review & editing, Software, Methodology, Formal analysis. Leslie Arkless: Writing – review & editing, Methodology, Formal analysis. Nina Cardinal: Writing – review & editing, Funding acquisition, Conceptualization. Maciej Zajac: Writing – review & editing, Methodology, Investigation, Conceptualization.

#### Declaration of competing interest

The authors have no competing interests to declare.

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#### Data availability

Data will be made available on request.

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