Case study

Assessment of activity moduli and acidic resistance of slag-based geopolymer concrete incorporating pozzolan

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The environmental impact of Portland cement production and utilization in the construction sector has led to the global call for the use of eco-friendly construction materials for the production of cleaner and sustainable products. Therefore, this study explored agro-industrial wastes, slag and corncob ash, for the production of geopolymer concrete (GPC). Corncob was dehydroxylated at 600°C for 3 h and partially used as a replacement for slag at 0%, 20%, 40%, 60%, 80%, and 100%. A 12 M, 14 M, and 16 M of both sodium silicate (SS) and sodium hydroxide (SH) were used as activators. The chemical moduli of each and mixed binder were quantified and evaluated based on the major reactive oxides, hence leading to the evaluation of reactivity indexes (RIs). Moreover, the RIs and mix design properties (MDPs) of concrete were used for the prediction of flexural strength while the chemical resistance of each concrete sample was investigated. Compared with the experimental results, the predictive flexural strengths based on the RIs and the MDPs yielded a high precision with \(R^2\) ranging from 88–92% at 7–90 days, respectively. Moreover, the GPC, unlike Portland cement concrete (PCC), resisted the more acidic attack. Therefore, the use of GGBFS–CCA blended concrete would be more advantageous in a highly acidic environment than PCC. Ultimately, the models proposed by this study can be useful in the concrete mix design procedure for the flexural strength development of GPC incorporating agro-industrial provided the oxide compositions of each and mixed material were obtained.

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1. Introduction

In the construction sector, the utilized rate of concrete is high, owing to the rapid industrialization and urbanization [1]. Amongst the concrete constituents, Portland cement (PC) plays a pivotal role in determining the quality of concrete. However, the production of PC, apart from its negative impact on the environment, requires a massive industrial process [2]. A ton of PC production, which emits 1 ton of carbon dioxide (\(\text{CO}_2\)) into the atmosphere, requires 4000 MJ energy, 1.5 tons of raw materials, and 140 kWh of electricity [3]. Moreover, a 7–9% of \(\text{CO}_2\) is emitted yearly into the atmosphere following the massive requirements of energy in PC production, hence contributing to the serious global warming [4]; this poses huge threats to human and ecosystem survival and development. Besides, from the building activities alone, Mahmoudkelaye et al. [5] estimated 30–40% generation of greenhouse gas (GHG) emissions, globally. Moreover, the yearly utilization of PCC in the construction industry is estimated to be 20 billion tons globally [6]. Furthermore, the need for the construction of
infrastructures in fast-growing cities could emit 226 gigatones of CO₂ by 2050 in the developing nations [6]. Following this trend, the carbon budget of 800 gigatones of total CO₂ emissions targeted by the Paris Climate Agreement after 2017 would be challenging to achieve.

In the construction sector, such as concrete mix design [7], pavement engineering [8], and geotechnical engineering [9], sustainable development and production have been a major priority. In the construction sector, there is a global call for the reduction of CO₂ emissions associated with PC production. The utilization of supplementary cementitious materials (SCMs) such as fly ash [1,10], coal ash [11], silica fume [1,12], metakaolin (MK) [10] ground granulated blast furnace slag (GGBFS) [10,12], rice husk ash (RHA) [13,14], corn cob ash (CCA) [15–17], cashew nutshell ash (CNSA) [18], bagasse ash (BA) [11], palm oil fuel ash (POFA) [19], and cassava peel ash (CPA) [20] in the production of sustainable construction materials, have been yielding favourable results in the recent studies. The partial or full incorporation of SCMs for the green concrete production, apart from limiting the environmental impact of PCC, has been reported to improve the workability [21,22]; mechanical [23,24]; durability [23,25]; and economical [26,27] properties. Moreover, the use of SCMs does not only reduce the initial CO₂ emissions of PC production but also reduce the volume of PCC needed in the construction industry and extend the building’s service life [4]. It is interesting to note that in 2016 and 2017 alone, the global production of corn was 969.69 and 1071.51 million metric tons, respectively [28]. Notwithstanding, most of the corn cobs produced are discarded as waste, hence culminating in environmental pollution; this justified the recycling of CCA for PC production. Besides, CCA possesses high silica and alumina contents, hence contributing to its strong pozzolanic response [15,21].

Different techniques have been used to correlate both mechanical and mix properties of concrete. Hammoudi et al. [29] applied artificial neural networks (ANNs) and response surface methodology (RSM) to evaluate the compressive strength (fc) of concrete. It was discovered that ANNs yielded better accuracy than RSM. Moreover, Al-Shamiri et al. [30] adopted an extreme learning approach (ELA) to predict the fc of high-strength concrete. It was revealed that ELA yielded acceptable precision to correlate the fc of high-strength concrete and the mix design proportions. Sobhani et al. [31] used the regression models to predict the fc of no-slump concrete and compared with ANNs and adaptive network-based fuzzy inference systems (ANFIS). It was inferred from the findings of the study that ANNs and ANFIS yielded better accuracy than regression models. However, the regression models offered a predictive equation for fc based on mix design proportion, while ANNs and ANFIS could not offer such an equation. Oyebisi et al. [18] and Xie and Visintin [14] applied the reactivity, hydraulic, and lime moduli to predict the relationship between the fc of concrete blended with SCMs and the mix design proportions such as water-to-binder ratio and binder-to-aggregate volume ratio of 0.618 and 0.30–1.5, and 0.12 and 0.045 to 0.359, respectively; the predictions yielded good accuracy with 99.66 % and 83.70 % coefficients of determination, respectively. Furthermore, the activity index of any SCM is influenced by its oxide composition, mineralogical composition, fineness, and specific surface area [32,33]. Besides, type, mineralogical, and chemical compositions of aggregates influence the performance and reactivity of concrete [34,35]. However, a single oxide cannot be used to quantify the reactivity of SCM. Hence reactivity, hydraulic, and lime moduli are majorly applied to quantify the hydraulic or self-cementitious properties, while both silica and alumina moduli are commonly used to determine pozzolanic properties [36,37].

Several standards have established the procedures of assessing the chemical indexes and hydraulic efficiency index of GGBFS [38,39] and the pozzolanic activity of pulverized fly ash or natural pozzolan [40,41]. ASTM C 989 [39] classifies GGBFS into three grades (80, 100, and 120) based on the mortar strength of slag activity index (SAI). The SAI for grade 80, 100, and 120, using 50 % cement replacement by the mass of the binding materials at both 7 days and 28 days must be 70 % and 80 %, 70 % and 90 %, and 90 % and 110 % strength minimum of the reference-cement mortar, respectively. Moreover, based on the caustic soda test in assessing the hydraulic activity of slag, ASTM C 1073 – 18 [42] recommends the compressive strengths of 7 MPa and 8 MPa, and 12 MPa and 15 MPa after 6 h and 8 h hardening, respectively. On the other hand, BS 3892 – 1 [40] specifies a SAI greater than 0.80 as a positive pozzolanic activity for fly ash or natural pozzolan for 30 % cement replacement after 7 days and 28 days. In contrast, ASTM C 618 [43] recommends a SAI greater than 0.75 for 20 % cement replacement after 7 days and 28 days. Despite the satisfactory and positive results of using chemical indexes and established procedure in assessing both hydraulic and pozzolanic properties of SCMs, there was no literature related to the activity indexes and durability properties of GPC incorporating GGBFS and CCA.

Many of the previous studies considered fc as the most remarkable factor to determine the quality of the concrete mixture [30,31]. Still, another factor, such as flexural strength (fr) has not been considered. Therefore, this paper provides new insight into the oxide compositions of each binding material by using the x-ray fluorescence analyzer (XRF); this guides in assessing the hydraulic and pozzolanic activities of GPC blended with agro-industrial wastes using the RI’s concept. It also provides an evaluation of the reactivity of each and blended binding material using the existing RIs, which are commonly used in self-cementitious and pozzolanic reactions; it develops a model to predict the fr of GPC blended with SCMs following the RI and mix design proportions, and investigates the performance of concrete produced under acidic attacks. In achieving these objectives, agro-industrial wastes such as GGBFS and CCA were harnessed, recycled, activated with alkaline solutions, and cured under ambient conditions. The experimental data for flexural strengths were obtained through laboratory work, while the predictive flexural strength was developed via the fit regression model in Minitab 17 to offer a predictive equation [31]. The hydraulic and pozzolanic activity tests of the binding materials were also examined using slag activity index and CST, and SAI and FT, respectively. Slag activity index and CST were adopted because they have been standardized and offered satisfactory results [37–39,42]. In the same vein, both SAI and FT have been standardized, reported, and significantly correlated [37], hence justifying their adoption in this present study. The recycling of both GGBFS and CCA would lessen the environmental, economic, and societal threats posed by the PC production; improve the concrete properties, and reduce the
construction cost and solid wastes, hence driving sustainability. The models developed from this study would enhance the findings of future studies on GPC incorporating SCMs by providing means of predicting fracture based on RIs and MDPs.

2. Materials and methods

2.1. Materials

The locally sourced materials, GGBFS and CCA, as shown in Fig. 1, were used as SCMs for the production of GPC, while Portland limestone cement (PLC), as shown in Fig. 1, was used as a binder for the production of PCC and compared with GPC. Slag was ground to obtain GGBFS. Corn cob was dehydroxylated at 600 °C for 3 h to obtain CCA. The SCMs were then sieved with BS 90 μm to obtain a similar fineness with PLC.

The specific gravity (SG) of the binding materials was determined following the requirements stated in BS EN 196–3 [44] using a specific gravity bottle and kerosene. The results indicated 2.90 g/cm³, 2.44 g/cm³, and 3.15 g/cm³ for GGBFS, CCA, and PLC, respectively. Owing to these results, GGBFS met the SG limit of 2.90 g/cm³ to 3.15 g/cm³ specified by BS EN 15167–1 [38], while that of CCA confirmed the similar results obtained by Oyebisi et al. [21].

The fineness of binding materials was determined using the dry sieving method, and BS sieve 90 μm as stipulated by BS EN 196–6 [45]. The results showed 7.6 %, 8.0 %, and 7.5 % for GGBFS, CCA, and PLC, respectively, hence satisfying the 12 % maximum fineness specification prescribed by BS EN 196–6 [45]. Therefore, the materials are suitable for use as binder and SCMs in concrete production. Furthermore, Laser diffraction, Model Beckman Coulter LS-100, was used to analyze the particle size distribution of the binding materials, as shown in Fig. 2, over the range size of 0.5 μm–900 μm. The results indicated a mean particle size of 20.68 μm, 23.45 μm, and 18.79 μm for GGBFS, CCA, and PLC, respectively. Besides, the specific surface area was carried out on the binding materials following the procedure stated by BS EN 196–6 [45] using the Blaine method at a standard porosity of 0.500. The results indicated 420 m²/kg, 625 m²/kg, and 375 m²/kg for GGBFS, CCA, and PLC, respectively.

The alkaline solutions, SH pellets with 99 % purity, and SS gel were locally sourced and used as activators. SS gel comprises Na₂O, SiO₂, and H₂O of 9.4 %, 30.1 %, and 60.5 % respectively, with SiO₂/Na₂O weight ratio of 3.20 and S.G. of 1.40 g/cm³ at 20 °C. A 354 g, 400 g, and 443 g of SH pellets were measured and dissolved in 646 g, 600 g, and 557 g of clean water based on the chemistry procedures established by Rajamane and Jeyalakshmi [46] for the preparation of 12 M, 14 M, and 16 M activators, respectively. The SH solutions were prepared 24 h earlier to reduce the high rise in temperature owing to the reaction between SH pellets and water and added to SS gel 2 h before casting for better performance, using a SS/SH ratio of 2.5:1.

The locally sourced aggregates were used and prepared at saturated surface conditions before the mix design. Grading was also conducted on the aggregates to obtain the needed particle size distribution (PSD). Moreover, the aggregates were characterized in line with the BS EN 12,620 [47]. The specific gravity (SG), water absorption (WA) and moisture content (MC) of the aggregates were determined following the procedure stated in BS EN 12,620 [47]. The results showed the SG of 2.60 g/cm³ and 2.64 g/cm³; WA of 0.7 % and 0.8 %, and MC of 0.3 % and 0.2 % for both fine aggregate (FA) and coarse aggregate (CA), respectively. Fig. 3 shows the PSD of both FA and CA used; the aggregates satisfied the limits of BS EN 12,620 [47]. On the other hand, the mineralogical composition of the coarse aggregate (granite) was identified with the aid of the Petrological Microscope, Model RPI-3 T. The sample was prepared, polished in a glass ground plate using a carborundum, and mounted on a clean glass slide with adhesive [48]. Also, the chemical composition was analyzed using the XRF spectrometer machine, Philips PW-1800. The results of mineralogical composition showed quartz, feldspar, mica, and iron oxide of 62.50 %, 20.45 %, 16.55 %, and 0.50 %, respectively. Moreover, the chemical composition reveals SiO₂, Al₂O₃, Fe₂O₃, CaO, MgO, SO₃, K₂O, Na₂O, P₂O₅, MnO, and LOI as 67.05 %, 14.40 %, 5.63 %, 3.90 %, 1.72 %, 0.02 %, 5.50 %, 1.16 %, 0.15 %, 0.05 %, and 0.52 %, respectively. From these results, it is inferred that the coarse aggregate is acidic granite because the content of SiO₂ was in the range of 66–75% [35]. Besides, based on alkalinity, the granite was classified as calcalkalinity in that (Na₂O + K₂O)/SiO₂ = 43 was 1.85; this

![Fig. 1. Binding materials used (a) PLC (b) GGBFS (c) CCA.](image-url)
ranged between 1.2–3.5 for calcalkalinity [35]. XRF analysis was not performed on the FA because it comprises SiO₂ content, almost in its entirety [48].

2.2. Tests for pozzolanic activity

2.2.1. Strength activity index (SAI)

The SAI was determined in line with the BS 3892–1 [40]. The water to binder ratio was modified to allow for the same flow properties with reference-cement mortar cubes [49], hence requiring mixing water of 235 mL of distilled water. Therefore, the SAI was determined for both 7 days and 28 days compressive strengths (CS) on the average of three samples using the relationship, as illustrated in Eq. 1 [40].

\[
\text{SAI, } \% = \frac{P}{C} \times 100
\]

where \( P \) is the average CS of pozzolan-reference cement mortar cubes (in MPa)

\( C \) is the average CS of reference-cement mortar cubes (in MPa).

The cement-reference mortar cubes, at 7 days and 28 days, showed the CS of 40.64 MPa and 50.43 MPa, respectively. Following the formula, as illustrated in Eq. 1, the test pozzolan (CCA) exhibited a SAI of 0.85 and 0.91 at 7 days and 28 days, respectively, thus showing considerable pozzolanic activity because SAI is greater than 0.80, as recommended by BS 3892–1 [40].
2.2.2. Frattini test (FT)

The FT was analyzed following the procedure specified by BS EN 195–5 [50]. The theoretical maximum concentration (TMC) of [CaO] was determined using the relationship, as illustrated in Eq. 2 [50]. As shown in Table 1, the [CaO] was compared with TMC [CaO], and the result was determined as the difference between the two values, expressing as a percentage of TMC removed. The pozzolanic activity of CCA, as shown in Table 1, removed the lime with 51 %, thus indicating that CCA is pozzolanic active.

$$\text{TMC} \{\text{CaO}\} = \frac{350}{[\text{OH}]} - 15$$

2.3. Tests for hydraulic activity

2.3.1. Slag activity index

The method was carried out following the procedure in ASTM C 989 [39] for the GGBFS. The procedure is similar to that of SAI stated in BS 3892–1 [40] except for 50 % cement replacement, C₃A content ranging from 6 to 10 %, and a maximum of 3% SO₃ content specified by the standards for the reference cement. Therefore, from the XRF results, PLC exhibits 2.03 % SO₃ content, hence satisfying the maximum requirement of 3%. Besides, C₃A was quantified based on Bogue’s equation, as shown in Eq. 3 [34]. Based on the XRF result and in line with Eq. 3, the result exhibited 10 % C₃A, thus fulfilling the maximum specification of 10 %. Finally, the slag activity index was determined following the relationship, as illustrated in Eq. 1.

$$\text{C₃A} = 2.65(\text{Al}_{2}O_{3}) - 1.69(\text{Fe}_{2}O_{3})$$

The CS of GGBFS-reference cement mortars with the mean particle size (d₅₀ = 20.68 μm) exhibited the slag activity index of 76.42 % and 98.53 % at 7 days and 28 days, respectively, hence classifying as grade 100 because the activity index is more than 70 % and 90 % at 7 days and 28 days, respectively [39].

2.3.2. Caustic soda test (CST)

This method was carried out following the procedure outlined in ASTM C 1073–18 [42]. The diluted solution-to-GGBFS ratio was fixed at 0.5 and used to prepare 40 mm × 40 mm × 160 mm prismatic samples. After 6 h and 24 h, all samples were demoulded and tested for CS. The results, average of three test samples, indicated 7.63 MPa and 14.52 MPa at 6 h and 8 h, respectively, hence satisfying the specifications of 7 MPa to 8 MPa and 12 MPa to 15 MPa after 6 h and 8 h, respectively, as recommended by ASTM C 1073–18 [42].

2.4. Materials characterization

The oxide compositions of binding materials, CCA, GGBFS, and PLC, were analyzed using the XRF spectrophotometer machine, Philips PW-1800. The results are shown in Fig. 4. The results revealed that CCA satisfied the chemical pozzolanic requirements stipulated by BS EN 450–1 [51] and BS EN 8615–2 [52] such that the addition of SiO₂, Al₂O₃, and Fe₂O₃ met 70 % minimum requirement. The content of CaO within the range of 10–20% established by Al-Akrhas [53] was also met. It can be deduced that the CCA could exhibit a pozzolanic reaction and used as the SCM in the production of blended GPC. On the other hand, GGBFS met the BS EN 15167–1 [38]’s limit requirements of 32–40% for both silica (SiO₂) and lime (CaO) contents. Besides, (\text{CaO} + \text{MgO}/\text{SiO}_2) ≥ 1, (\text{CaO}/\text{SiO}_2) ≤ 1.4, and SiO₂ + CaO + MgO ≥ 67 % stipulated by BS EN 15167–1 [38] were also met. Also, the oxide compositions obtained herein for GGBFS showed similar compositions with the previous studies [19,31]. Therefore, an inference is made that GGBFS utilized in this study could exhibit both pozzolanic and self-cementitious reactivity, hence suitable for use. In the same vein, the PLC fulfilled the chemical requirements specified by BS EN 196–2 [54].

The microstructural behaviour of the binding materials, GGBFS, CCA, and PLC, was examined using the SEM machine, JEOL 7000600, to establish the characteristics that influenced the RIs of each binder. The SEM analysis was performed on a flat (general) scan. For the investigation, the accelerated voltage was constant at 15 kV, while images were observed at 4000x magnification in a high vacuum. The SEM micrograph results are presented in Fig. 5(a), to a limited extent, reveals a wrinkled internal structure with sharp needles. However, Fig. 5(b) shows an amorphous structure, while Fig. 5 (c) reveals a crystalline and spherical structure.

Table 1

| Sample | [OH] mmol l⁻¹ | [CaO] mmol l⁻¹ | TMC [CaO] mmol l⁻¹ | [CaO] reduction (%) |
|--------|---------------|----------------|-------------------|---------------------|
| Control | 56.83         | 8.25           | 8.37              | 0.70               |
| CCA    | 39.72         | 4.63           | 14.16             | 50.72              |
2.5. Mix design quantities

The mix quantities were designed following the procedures stated by BS EN 206 [55]. The percentage replacement of GGBFS by CCA was selected to examine the replacement levels, which would meet the target strengths for both structural and non-load bearing applications. Owing to this, GGBFS was replaced with CCA at 0%, 20%, 40%, 60%, 80%, and 100% for the production of GPC and was respectively indicated as E1, E2, E3, E4, E5, and E6, while the PLC (100% GGBFS) was indicated as E0. The mix was designed to attain target strengths 30 MPa and 40 MPa for grades M30 and M40 concrete, respectively. The mix design quantities for both M30 and M40 are shown in Tables 2 and 3, respectively.

2.6. Mix preparation, casting and curing

The dry constituents were prepared following the procedures prescribed by BS 1881–125 [56] and BS EN 12390–2 [57] by preparing and pouring fresh concrete into a cubical mould of 150 mm x 150 mm x 150 mm long beam for flexural strength test. The fresh sample was randomly compacted each in three layers, cured under 25℃ and 65% RH. The compressive strength was tested at 90 days curing, while the flexural strength was tested at 7, 28, 56, and 90 days.
2.7. Experimental tests and analysis

2.7.1. Mechanical test

The compressive strength (fc) and flexural strength (fr) were determined with the aid of an INSTRON 5000R UTM following the procedures stated by BS EN 12390–4 [58] and BS EN 12390–5 [59] in a constant force regime under a loading rate of 0.6 MPa and 0.06 MPa per second, respectively. Three (3) samples were made and crushed for each mix ID, and the average was used for the analysis.

2.7.2. Reactivity indexes (RIs) of binding materials

The RIs of binding materials were evaluated using the principal reactive oxides such as CaO, SiO₂, Al₂O₃, Fe₂O₃, MgO, and SO₃ following the establishment of their oxide compositions, which reflect both self-cementitious and pozzolanic reactivity [14,18,32]. The concept which guides the RIs is illustrated in Eq. 4–8 as reactivity, hydraulic, lime, silica, and alumina moduli of each and blended binder, indicating as RM, HM, LM, SM, and AM, respectively.

\[
RM = \frac{\text{CaO} + \text{MgO} + \text{Al}_2\text{O}_3}{\text{SiO}_2}
\]  

(4)

\[
HM = \frac{\text{CaO}}{\text{SiO}_2 + \text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3}
\]  

(5)

\[
LM = \frac{1.0\text{CaO} - 0.7\text{SO}_3}{2.8\text{SiO}_2 + 1.1\text{Al}_2\text{O}_3 + 0.7\text{Fe}_2\text{O}_3}
\]  

(6)

\[
SM = \frac{\text{SiO}_2}{\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3}
\]  

(7)

\[
AM = \frac{\text{Al}_2\text{O}_3}{\text{Fe}_2\text{O}_3}
\]  

(8)

Table 2
Mix quantities for M 30 (in Kg/m³).

| Mix ID | PLC | GGBFS | CCA | FA | CA | SH | SS | SS/SH |
|--------|-----|-------|-----|----|----|----|----|-------|
| E0     | 390 | 0     | 0   | 675| 1031| 0  | 0  | 0     |
| E1     | 0   | 390   | 0   | 675| 1031| 0  | 0  | 0     |
| E2     | 0   | 312   | 78  | 675| 1031| 0  | 0  | 0     |
| E3     | 0   | 234   | 156 | 675| 1031| 0  | 0  | 0     |
| E4     | 0   | 156   | 234 | 675| 1031| 0  | 0  | 0     |
| E5     | 0   | 78    | 312 | 675| 1031| 0  | 0  | 0     |
| E6     | 0   | 0     | 390 | 675| 1031| 0  | 0  | 0     |

w/b is the water to binder ratio = 0.42; b/agg is the binder to aggregate ratio = 0.31.

Table 3
Mix quantities for M 40 (in Kg/m³).

| Mix ID | PLC | GGBFS | CCA | FA | CA | SH | SS | SS/SH |
|--------|-----|-------|-----|----|----|----|----|-------|
| E0     | 500 | 0     | 0   | 585| 1031| 0  | 0  | 0     |
| E1     | 0   | 500   | 0   | 585| 1031| 0  | 0  | 0     |
| E2     | 0   | 400   | 100 | 585| 1031| 0  | 0  | 0     |
| E3     | 0   | 300   | 200 | 585| 1031| 0  | 0  | 0     |
| E4     | 0   | 200   | 300 | 585| 1031| 0  | 0  | 0     |
| E5     | 0   | 100   | 400 | 585| 1031| 0  | 0  | 0     |
| E6     | 0   | 0     | 500 | 585| 1031| 0  | 0  | 0     |

w/b is the water to binder ratio = 0.54; b/agg is the binder to aggregate ratio = 0.23.
2.7.3. Prediction of $f_r$ based on RIs and MDPs

For this study, MDPs are indicated as water to binder (w/b) ratio and binder to aggregate (b/agg) ratio. Either reactivity, hydraulic, or lime moduli quantifies the self-cementitious properties of each and blended binding material while the pozzolanic activity is quantified by both silica and alumina moduli [32,37]. Consequently, a linear relationship exists in the prediction of $f_r$ and RIs. Thus, the regression was first modelled based on the combination of RM, SM, and AM; HM, SM, and AM; and LM, SM, and AM using Minitab 17 statistical software. Furthermore, in determining the $f_r$ of blended concrete, the RIs of blended binders were integrated and normalized with an inverse of w/b ratio; hence, $f_r$ becomes a direct proportion to RIs, but an inverse proportion to w/b ratio [14,18]. Therefore, the fit regression relationship between $f_r$ and w/b ratio was first normalized and modelled in the range of 0.54 to 0.42 w/b ratio for grades M 30 to M 40 concrete, respectively. The $f_r$ and RIs were selected as the response (dependent variable) and continuous predictors (independent variables), respectively, to predict the design data in Minitab 17.

The binder to aggregate (b/agg) ratio also contributed a vital role to the evaluation and improvement of the concrete strength apart from RIs and w/b ratio [14]. The $f_r$ of blended binders was significantly improved when RIs, w/b ratio, and b/agg were all used for the strength correlation. It is noteworthy to state that the volume ratio was used to model the b/agg ratio against the weight ratio. For each mix, the volume fraction was determined using its moisture content and specific gravity to improve the binder-aggregate packing capacity. Following the incorporation of w/b ratio, the fit regression relationship between $f_r$ and b/agg ratio was modelled in the range of 0.31 to 0.23 b/agg ratio for grades M 30 to M 40 concrete, respectively. Consequently, $f_r$ was predicted based on the RIs and MDPs, as illustrated in Eq. 9–11.

$$f_r = \beta + \left(\frac{\alpha_1 RM + \alpha_2 SM + \alpha_3 AM}{w/b}\right)(b/agg)$$  \hspace{1cm} (9)

$$f_r = \beta + \left(\frac{\alpha_1 HM + \alpha_2 SM + \alpha_3 AM}{w/b}\right)(b/agg)$$  \hspace{1cm} (10)

$$f_r = \beta + \left(\frac{\alpha_1 LM + \alpha_2 SM + \alpha_3 AM}{w/b}\right)(b/agg)$$  \hspace{1cm} (11)

where $\beta$, $\alpha_1$, $\alpha_2$, $\alpha_3$ represent the magnitudes of coefficients.

2.7.4. Durability test

The chemical resistance was conducted on the prepared cube samples using the solutions of sulphuric acid (H₂SO₄) at 2% concentration [34,60] for acidic attacks. The concrete specimens were tested for both weight and strength loss after 90 days of immersion in H₂SO₄.

![Graphs showing compressive strength for different mixes](image)

**Fig. 6.** Compressive strength (a) M 30 (b) M 40.
3. Results and discussion

3.1. Mechanical properties

Figs. 6 and 7 present the results of both compressive and flexural strengths, respectively. From Figs. 6 and 7, the results revealed both compressive and flexural strengths increased with increasing GGBFS content for both M 30 and M 40 at all curing days, respectively. The results could be associated with the reaction between the aluminosilicate glassy phases (amorphous structure) of GGBFS, as shown in Fig. 5(b), and the alkaline solutions, hence resulting in x-ray amorphous aluminosilicate paste (X-RAAP). The X-RAAP, according to Criado et al. [61], contributes to the higher strengths of the hardened product, compared with both wrinkled and crystalline structures for PCC and CCA, as shown in Fig. 5(a) and (c), respectively. Besides, unlike 12M and 16M activators, 14M activator exhibited the highest compressive and flexural strengths at all curing days because of its capacity to liberate more aluminosilicate gels in the mix. However, at 16M activating level, the OH⁻ solution in the mix could be excessive, thus encasing the amorphous paste, causing a barrier to the activating dissolution, and delaying the hydrating agent (calcium-silicate-aluminate-hydrate, C-S-A-H) in the mixed paste; this delays and decreases the strengths.
3.2. Principal reactive oxides of blended binders

Fig. 8 shows a decrease in CaO, Al₂O₃, and MgO with increasing CCA content, while SiO₂, Fe₂O₃, and SO₃ increase with increasing CCA content in the blended mix; this supports the similar findings reported by Akinwumi and Aidomoje [17] that the reactive oxides, CaO, MgO, and Al₂O₃ decrease with increasing CCA content, while SiO₂, Fe₂O₃, and SO₃ increase with increasing CCA content for the CCA–PC blend. Meanwhile, Behim et al. [32] and Demoulian et al. [36] stated that GGBFS exhibits a similar mineralogical composition to PC; it majorly possesses oxides of Ca, Si, Al, Mg, and Fe, and this gives GGBFS its hydraulic and pozzolanic properties. Also, Xia and Visintin [14] and Darquennes [33] opined that slag is said to exhibit both self-cementitious and pozzolanic properties if the content of CaO and SiO₂ is higher than 30 %. From the XRF results of GGBFS, it is clear that the contents of both CaO and SiO₂ are higher than 30 %. On the other hand, Taylor [62] and Hewlett [63] reported that the self–cementitious reaction of slag decreases as the crystalline content in the blended mix increases; this demonstrates that the reactivity of GGBFS depends on the increasing content of its amorphous structure, and the significant oxides which contribute to the high phase of an amorphous structure are oxides of Ca, Al, and Mg [64,65]. Thus, through close examination of microstructures of binding materials, as shown in Fig. 5, it was evident that the content of the amorphous structure in GGBFS could gradually decrease while the content of the crystalline structure in CCA might increase when GGBFS is replaced with CCA. Consequently, as the content of CCA in the blended mix increases, CaO, Al₂O₃, and MgO in GGBFS decrease. In contrast, SiO₂, Fe₂O₃, and SO₃ in CCA increase; this corroborates the findings from relevant studies in that the reactivity of GGBFS increases with increasing CaO, MgO, and Al₂O₃ contents but reduces as the contents of SiO₂, Fe₂O₃, and SO₃ rise [66,67]. However, it was pointed out that GGBFS comprises small crystal material and is advantageous to its reactivity [68–70]. Besides, Gruskovanjak et al. [67] pointed out that the optimum content of the principal reactive oxides of slag is more beneficial to its self-cementitious reactivity than the content of the amorphous structure. Therefore, it is inferred that the contents of CaO, Al₂O₃, MgO, SiO₂, Fe₂O₃, and SO₃ influence the reactive potentials of GGBFS–CCA blended binders.

3.3. RIs of the blended mix

In assessing the RIs of each blended binder, Eq. 4–5 was used, and the results are shown in Fig. 9. It was revealed that the RM, HM, LM, and AM decreased with increasing CCA content, while the SM increased with increasing CCA content in the blended mix for both M 30 and M 40. Besides, it was evident from the results that CaO, Al₂O₃, MgO, SiO₂, Fe₂O₃, and SO₃ influenced the RIs of the blended binders. The RM, HM, and LM of the blended binders increased with increasing CaO, Al₂O₃, MgO contents, while the SM and AM of the blended binders increased as the contents of SiO₂ and Al₂O₃ increased, respectively. In contrast to HM, the RM of the blended binders met the minimum requirement of 1.0 specified by BS EN 8615–2 [52]. Statistically, the RM, HM, LM, and AM of the blended mix decreased from 25 to 78 %, 19–77%, 19–77%, and 11–26% as the percentage replacement of CCA by GGBFS increased from 20 % to 100 % for both M 30 and M 40, respectively. The self-cementitious properties of mixed binders increase with an increase in CaO, Al₂O₃, and MgO contents, thus resulting in stronger hydraulic reactions [32,71]. As a result, the decrease in RIs may be attributed to the reduction in principal reactive oxides, CaO, Al₂O₃, and MgO as a result of the increase in CCA content in the blended mix. However, the SM of the blended mix decreased from 44 to 10 % as the percentage replacement of CCA by GGBFS rose from 20 to 100 % for both M 30 and M 40, respectively. Meanwhile, it is shown in Fig. 7 that CCA, being a pozzolan, exhibits higher content of silica (SiO₂) compared with that of GGBFS. Therefore, this result confirms the findings reported by Mathhes et al. [70] that SM increases with increasing SiO₂ content, hence resulting in stronger pozzolanic properties. On the other hand, the reactivity of

![Fig. 9. RIs of each and mixed binder for (a) M 30 and (b) M 40.](image)
GGBFS depends on its amorphous structure, thus influencing its RIs [69,70]. This assertion confirms the SEM micrographs, as shown in Fig. 5 (b) and (c) for GGBFS and CCA, which display amorphous and crystalline structures, respectively.

3.4. Prediction of $f_c$ based on RIs and MDPs

3.4.1. Prediction of $f_c$ based on RM, AM, SM, and MDPs

Following Eq. 9, the results of the statistical data are shown in Fig. 10 (a)–(d) for 7, 28, 56, and 90 days, respectively. It was observed that some data points for SM significantly deviated from the regression line. This may be asserted to the diversity of chemical compositions of blended binders, aggregate type and volume, and mix design proportions; this assertion confirms the findings reported by Xie and Visintin [21] and Neville [34] that differences in the oxide composition of blended binders, aggregate types, texture, and shape, and methods of mix design, affect the data results, hence influencing the reactive potentials of blended concrete incorporating SCMs. Moreover, the flexural strength increased with increasing RM and AM but decreasing SM; this may be attributed to the higher contents of CaO and Al₂O₃ in GGBFS, which increases RM and AM, thus resulting in a stronger self-cementitious reaction. However, the higher content of SiO₂ in CCA increases its SM, hence leading to a pozzolanic reaction rather than a hydraulic reaction; this also confirms the findings of a similar study reported by Gruskovnjak et al. [67] that the RM and AM increase as the CaO and Al₂O₃ contents increase, while SiO₂ content reduces, thus resulting in high reactivity. However, the higher contents of SiO₂ and low contents of CaO and Al₂O₃ result in low reactivity. On the other hand, a blended mix with high contents of CaO, Al₂O₃, and MgO exhibits high self-cementitious/hydraulic properties in the presence of alkaline activators [66,68,70].

The fit regression model was used for the correlation of $f_c$ based on the RIs (RM, AM, and SM) and MDPs at the global trend of 95 % confidence interval (CI) and prediction interval (PI). Thus, the regression equations are illustrated in Eq. 12–15 for 7, 28, 56, and 90 days, respectively. Therefore, the coefficient of determination ($R^2$) is 87.47 %, 87.60 %, 88.12 %, and 92.26 % fit to predict the data at 95 % CI and PI for 7, 28, 56, and 90 days, respectively, thus
inducing 0–5.20% increase in $R^2$ as the curing age increases from 7–90 days. Ultimately, relative to RIs and MDPs, these developed models can be used for the strength prediction of concrete incorporating SCMs.

$$f_{r-7\text{days}} = \left( \frac{7.81\text{AM} - 1.468\text{SM} - 2.63\text{RM}}{w} \right) \left( \frac{b}{\text{agg}} \right) + 3.028$$  \hspace{1cm} (12)

$$f_{r-28\text{days}} = \left( \frac{2.22\text{AM} - 0.440\text{SM} + 1.66\text{RM}}{w} \right) \left( \frac{b}{\text{agg}} \right) + 3.825$$  \hspace{1cm} (13)

$$f_{r-56\text{days}} = \left( \frac{2.82\text{AM} - 0.5975\text{SM} + 1.10\text{RM}}{w} \right) \left( \frac{b}{\text{agg}} \right) + 4.125$$  \hspace{1cm} (14)

$$f_{r-90\text{days}} = \left( \frac{4.91\text{AM} - 0.969\text{SM} - 0.13\text{RM}}{w} \right) \left( \frac{b}{\text{agg}} \right) + 4.193$$  \hspace{1cm} (15)

3.4.2. Prediction of $f_r$ based on HM, AM, SM, and MDPs

Fig. 11 (a)-(d) indicates the statistical data for HM, AM, SM, and MDPs at 7, 28, 56, and 90 days, respectively. It was observed that some data points of SM were out of the regression line due to the difference in binders’ oxide compositions, the volume and chemical compositions of aggregates, and mix proportions. Besides, the $f_r$ of GGBFS–CCA blended concrete increased with increasing HM and AM but decreasing SM. The reason for a higher strength cannot be far-fetched: GGBFS exhibits higher content of CaO and Al$_2$O$_3$ compared with CCA, hence resulting in stronger hydraulic reaction, but this hydraulic reaction decreases when replaced with CCA, which predominantly contains a higher content of SiO$_2$. This assertion is in line with the findings reported in various studies that the hydraulic response of slag reduces with increasing silica content [64,65]. Therefore, it is inferred that the HM of GGBS–CCA blended binder increases with higher contents of CaO and Al$_2$O$_3$ and the lower content of SiO$_2$ in the mix.

![Graphs showing statistical data for HM, AM, SM, and MDPs](image)

**Fig. 11.** Statistical data for HM, AM, SM, and MDPs (a) 7 (b) 28 (c) 56 and (d) 90 days.
The fit regression model was used for the correlation of \( f_r \) based on the RIs (HM, AM, and SM) and MDPs at the 95 % CI and PI, and the regression equations are illustrated in Eq. 16–19 for 7, 28, 56, and 90 days. Thus, at 7, 28, 56 and 90 days, \( R^2 \) is 87.35 %, 87.84 %, 88.38 %, and 92.35 % fit to correlate the data, respectively. Statistically, there is 0–5.4% increase in \( R^2 \) as the curing age increases from 7 to 90 days.

\[
\begin{align*}
\frac{fr_{-7\text{ days}}}{} &= \left( \frac{5.50AM - 1.036SM - 1.99HM}{w} \right) \left( \frac{b}{agg} \right) + 3.022 \\
\frac{fr_{-28\text{ days}}}{} &= \left( \frac{2.60AM - 0.521SM + 2.82HM}{w} \right) \left( \frac{b}{agg} \right) + 3.814 \\
\frac{fr_{-56\text{ days}}}{} &= \left( \frac{2.59AM - 0.565SM + 2.59HM}{w} \right) \left( \frac{b}{agg} \right) + 4.111 \\
\frac{fr_{-90\text{ days}}}{} &= \left( \frac{3.69AM - 0.753SM + 1.49HM}{w} \right) \left( \frac{b}{agg} \right) + 4.178
\end{align*}
\]

3.4.3. Prediction of \( f_r \) based on LM, AM, SM, and MDPs

The statistical data for LM, AM, SM, and MDPs are indicated in Fig. 12 (a)-(d) for 7, 28, 56, and 90 days, respectively. It was noticed that some data points of SM were out of the global trend due to the diversity in oxide compositions, aggregates volumes and types, and mix proportions of the blended binders. Moreover, the \( f_r \) of GGBFS–CCA blended concrete increased with increasing LM and AM but decreasing SM; this may be attributed to the fact that GGBFS exhibits higher content of CaO and Al\(_2\)O\(_3\) compared with CCA, hence resulting in a stronger reactive component. Still, this reactive component decreases

Fig. 12. Statistical data for LM, AM, SM, and MDPs (a) 7 (b) 28 (c) 56 and (d) 90 days.
when replaced with CCA, which majorly contains a higher content of silica. Therefore, it is inferred that the LM of GGBFS–CCA blended binder increases with higher contents of CaO and Al₂O₃ and the lower content of SiO₂ in the mix.

The $f_r$, RIs (LM, AM, and SM), and MDPs were predicted using the fit regression model at the 95 % CI and PI, and the regression equations are illustrated in Eq. 20–23 for 7, 28, 56, and 90 days, respectively. Therefore, at 7, 28, 56, and 90 days, $R^2$ is 87.50 %, 87.65 %, 88.09 %, and 92.25 % fit to correlate data, respectively, hence indicating 0–5.20% increase in $R^2$ as the curing age increases from 7 to 90 days. Finally, with respect to RIs and MDPs, these proposed models can be used for the strength prediction of concrete incorporating SCMs.

$$f_{r-7\ days} = \left( \frac{6.51 \ AM - 1.250 \ SM - 7.74 \ LM}{W} \right) \left( \frac{b}{agg} \right) + 3.044$$ (20)

$$f_{r-28\ days} = \left( \frac{4.21 \ AM - 0.802 \ SM + 1.07 \ LM}{W} \right) \left( \frac{b}{agg} \right) + 3.835$$ (21)

$$f_{r-56\ days} = \left( \frac{3.87 \ AM - 0.786 \ SM + 1.62 \ LM}{W} \right) \left( \frac{b}{agg} \right) + 4.127$$ (22)

$$f_{r-90\ days} = \left( \frac{4.74 \ AM - 0.938 \ SM - 0.04 \ LM}{W} \right) \left( \frac{b}{agg} \right) + 4.192$$ (23)

### 3.4.4. Comparison of experimental results with predicted values

Fig. 13 illustrates the statistical comparison and trend between the flexural strengths of experimental results and that of predictive values. It was observed that both empirical and predictive results exhibited similar values and patterns of flexural strength. In contrast to HM, both LM and RM showed the best fit at all levels of curing time for both M 30 and M 40. These observations confirm the findings reported in similar studies such that LM yields the best fit for PC blended with cashew nut shell ash (CNSA) [18]. In contrast, RM yields the best fit for blended concrete incorporating SCMs [14]. Despite producing the
besides, variations and for against 2% GGBFS M blended binders, was low compared with the minimum requirements (≥ 0.66 ≤ 1.02) recommended by BS EN 197–1 [72]; besides, HM was less than 1 compared with the minimum requirement (< 1) specified by BS EN 197–1 [72]. However, RM satisfied the minimum requirement (< 1) defined by Behim et al. [32], Demoulian et al. [36], and BS EN 15167–1 [38]. The variations in LM and HM may be attributed to the difference in chemical and mix properties of concrete in that BS EN 197–1 [72]’s recommendation was based on the PC blended binders such that the ratio of CaO to SiO₂ in the blended mix was high compared with GGBFS–CCA blended binders reported in this study. Therefore, it is inferred that RM yields the best fit for GGBFS–CCA blended binder, and this can be used in the validation of blended binders incorporating SCMs.

3.5. Chemical attacks

Owing to the immersion of selected samples on 2% H₂SO₄ solution for 90 days, Fig. 14 (a) and (b) present the weight loss for both M 30 and M 40, respectively. While Fig. 15 (a) and (b) present the compressive strength (fₐ) loss for both M 30 and M 40, respectively. It was observed that the percentage weight loss in GPC samples, for both M 30 and M 40, ranged from 1 to 2% against 10–13% in PCC samples, while the percentage strength loss in GPC, for both M 30 and M 40, varying from 1 to 5% against 12–14% in PCC samples after 90 days exposure of concrete samples in 2% H₂SO₄. These results confirm the similar findings reported by Sanni and Khadiranaikar [25], Malhotra et al. [73], and Singh et al. [74] that the percentage weight loss and strength loss in GPC samples was less than 5% against 15% in PCC samples after 90 days of exposure in 2% H₂SO₄.

**Fig. 14.** Weight loss of cubes immersed in 2% H₂SO₄ for 90 days (a) M 30 (b) M 40.

**Fig. 15.** Strength loss of cubes immersed in 2% H₂SO₄ for 90 days (a) M 30 (b) M 40.
deterioration may be attributed to the aggressive attack of H2SO4 on Ca(OH)2 and C-S-H of PCC structure. In contrast, the C-S-A-H on GPC structure is protected and fixed by the presence of diluted water-glass (sodium silicate), thus forming calcium silicates which filled the pores [23,25,34]. Therefore, it is inferred that GGBFS–CCA blended GPC offers better resistance to acidic attack than PCC.

E1 (100 % GGBFS), E2 (80 % GGBFS + 20 % CCA), E3 (60 % GGBFS + 40 % CCA), E4 (40 % GGBFS + 60 % CCA), E5 (20 % GGBFS + 80 % CCA), and E6 (100 % CCA).

4. Conclusion

The study examined the GGBFS–CCA-based GPC, and its effects on the activity indexes and the acidic attacks were evaluated. Both experimental and statistical methods were used in the course of the study, and the results were compared with PCC. Consequent upon the findings and in line with research aims, the following sets of conclusions are made:

- The reactivity of GGBFS–CCA blended binder increases with increasing CaO, MgO, and Al2O3 contents. However, the reactivity decreases with increasing SiO2, Fe2O3, and SO3 contents.
- The RM, HM, and LM of GGBFS–CCA blended binder increases with increasing CaO, MgO, and Al2O3 contents, while the SM and AM increase with increasing SiO2 and Al2O3 contents.
- Flexural strength of GGBFS–CCA GPC increases with increasing RM, HM, LM, and AM.
- RM yields the best fit for predicting the flexural strength of slag-based GPC, incorporating CCA compared with HM and LM.
- Besides, a good correlation exists between the experimental results and proposed model equations.
- There is a remarkable improvement in R2 as the curing age increases.
- Slag-based GPC incorporating CCA provides excellent acidic resistance superior to that of PCC.

The concept of activity moduli in predicting the f_c of the GGBFS–CCA blended mix is attainable. This study benefits future research by focusing on three prospective solutions. First, the proposed model equations can be useful in the prediction and application of strength design proportions for GPC incorporating agro-industrial by-products under ambient curing conditions provided the chemical compositions are obtained. Second, the efficiency of the fit regression model in predicting f_c, based on the RIs and the MDPs is affirmed. Third, the application of agro-industrial by-products, GGBFS and CCA, can be advantageous in a highly acidic environment.

Declaration of Competing Interest

The authors show a credit to the sources in the manuscript. The authors declare that they have no known competing for financial interests or personal relationships that could have appeared to influence the work reported in this paper. The raw/processed data required to reproduce these findings cannot be shared at this time as the Data also forms part of an ongoing study. The authors declare that the manuscript is the authors’ original work and has not been published before. The authors also declare that the article contains no libellous or unlawful statements and does not infringe on the rights of others.

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