A New, Carbon-Negative Precipitated Calcium Carbonate Admixture (PCC-A) for Low Carbon Portland Cements

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Abstract: The production of Portland cement accounts for approximately 7% of global anthropogenic CO₂ emissions. Carbon CAPture and CONversion (CAPCON) technology under development by the authors allows for new methods to be developed to offset these emissions. Carbon-negative Precipitated Calcium Carbonate (PCC), produced from CO₂ emissions, can be used as a means of offsetting the carbon footprint of cement production while potentially providing benefits to cement hydration, workability, durability and strength. In this paper, we present preliminary test results obtained for the mechanical and chemical properties of a new class of PCC blended Portland cements. These initial findings have shown that these cements behave differently from commonly used Portland cement and Portland limestone cement, which have been well documented to improve workability and the rate of hydration. The strength of blended Portland cements incorporating carbon-negative PCC Admixture (PCC-A) has been found to exceed that of the reference baseline—Ordinary Portland Cement (OPC). The reduction of the cement clinker factor, when using carbon-negative PCC-A, and the observed increase in compressive strength and the associated reduction in member size can reduce the carbon footprint of blended Portland cements by more than 25%.

Keywords: CO₂ emissions; carbon CAPture and CONversion (CAPCON); Precipitated Calcium Carbonate (PCC); limestone; strength; rheology; XRD analysis; Life Cycle Assessment

1. Introduction

1.1. Background

Estimates vary, but the cement industry currently produces at least 2.8 billion tonnes of cement per annum, accounting for ~7% of global anthropogenic CO₂ emissions [1–4]. Cement production is expected to increase yearly to more than 4 billion tonnes per annum by 2040.

In 2009, the International Energy Agency (IEA) proposed the use of carbon capture and storage to reduce these emissions [5]. Since the IEA’s proposal to reduce the carbon emissions of the cement industry, new technologies have been developed to either capture and utilise post-combustion CO₂ or convert it into useful products. Some of these technologies are discussed in Section 1.2.

Calcium carbonate, in the form of limestone, has seen an increase in its use as a cement additive to reduce the carbon footprint of cement clinker, which is due to the relatively low CO₂ output of the process in comparison to Portland cement manufacture [6]. Most codes of practice allow a maximum of 5% ground limestone, containing >70% calcium carbonate, to be added to create Portland limestone cement (PLC) blends as distinct from Ordinary Portland Cement (OPC). Ground limestone has also been demonstrated to increase workability without impacting strength if used in quantities of less than or equal to the maximum allowable 5% in accredited PLC blends [7–9].
We present the first results of tests on the use of carbon-negative Precipitated Calcium Carbonate Admixture (PCC-A), a carbon CAPture and CONversion (CAPCON) product of the Carbon Capture Machine (CCM) (see http://www.ccmuk.com for more information), to reduce the carbon footprint of blended cements. Unlike ground limestone, which is carbon positive, PCC-A from the CCM CAPCON process is carbon negative and sequesters between 100 and 350 kgCO₂ per tonne of PCC-A produced, depending on the process inputs. Our proposal for a more sustainable future, where cement production is decarbonised at source, is illustrated in Figure 1.

![Figure 1. Proposal for the production of a new generation of enhanced performance, low carbon Precipitated Calcium Carbonate Admixture (PCC-A) blended Portland cements can be achieved in situ using the Carbon Capture Machine (CCM).](image)

The role of calcium carbonate in blended Portland cements spans chemical and physical properties. The chemical role is through the formation of (hemi- and mono-) carboaluminate phases during hydration [10]. Physically, it is predominantly an inert filler [8,11–13]. Unpublished data suggests that the control of grain size and the morphology of the PCC-A in the CCM’s CAPCON process can be beneficially used to produce blended Portland cements that do not compromise strength, durability or passivation while offering performance benefits that match or surpass OPC.

The test methods reported in this paper were used to evaluate, for the first time, the effects of adding carbon-negative PCC-A to OPC, potentially leading to the development of new classes of high performance, low carbon Portland cement blends. The methods were used to determine strength, rheology, passivation potential and the reduction in embodied CO₂ content. It has been shown that the use of PCC-A in blended cements can achieve an increased strength at contents well beyond the 5% limit for ground limestone, while concurrently reducing the embodied CO₂ content by 25% or more.

1.2. Carbon Capture and Utilisation in Cement Production

Portland cement is the most abundantly manufactured material globally and is also one of the largest sources of CO₂ emissions [1–4]; it is thus a prime candidate for the emerging carbon capture and utilisation industry to prioritise. A recent report published by Chatham House [14] estimates that, to effectively control the impact of the cement industry on the environment, the clinker factor (i.e., active fraction) in cements has to be reduced to 60% of the present value by 2050. Once this has been achieved, other methods have to be developed that rely on carbon capture and utilisation.

An alternative to carbon capture outlined in the Chatham House report is the use of “novel cements”—low carbon alternatives to traditional Portland cement blends. These cements include allegedly carbon-free magnesium oxide derived from magnesium silicates (MOMS) cements, calcium sulfoaluminate (CSA) cements and carbonated calcium silicate (CCSC) cements amongst others. However, novel cements have seen little market penetration due to their high production costs (MOMS
cements), specialty usage (CSA cements), scalability issues (CCSC cements) and resistance to change by codes of practice, consumer groups and standards institutes, compared to tried, tested and trusted Portland cement alternatives. The Chatham House report claims that carbon capture methods could reduce CO₂ emissions by 95–100%. Similarly, a study of several active carbon capture plants conducted by David and Herzog [15] found the typical efficiency to be 90% for direct emissions. In both cases, the conclusions relate to high cost, non-viable carbon capture and storage routes and are therefore of limited value. The estimates also ignore the contributions of secondary emissions from quarrying, transport and grinding, where the CO₂ cannot be easily captured—see Figure 2.

![Figure 2: Breakdown of CO₂ emissions of Portland cement production. Data taken from the Chatham House report [14].](image)

Worldwide, CO₂ capture and utilisation methods for the cement industry currently include the production of carbon-negative admixtures through mineralisation (Carbon Capture Machine) [6] and carbonation cured cements (Solidia, CarbonCure, CO₂NCRETE) [16–18].

Carbonation cured cements were first proposed in the 1970s, but their development has been stunted due to process complexity, cost and application scope. In the carbonation curing process, CO₂ is converted into the various polymorphs of calcium carbonate—calcite being the most common—through the interaction of CSH gel with CO₂ (gas) proceeding in two stages: initially the CSH “sheds” Ca forming CaCO₃ and a low Ca CSH limit is reached at Ca/Si ~1 which, upon continued carbonation, yields silica gel and calcium carbonate [19]. This differs from the normal hydration route where calcium hydroxide is formed in place of calcium carbonate [19]. Prior to CO₂ exposure, the free water content of the cement is reduced to ensure that the CO₂ can diffuse efficiently throughout. The cement is then exposed to CO₂ at pressures up to 5 atm. After exposure, the unreacted cement may be hydrated again to ensure that unreacted phases can continue to hydrate and provide further increases in strength [20]. The CO₂ content of carbonation cured cements was verified by Seo et al. [21] through XRD analysis and determined to range from 6.9 to 11% of the cement, depending on composition.

The carbonation curing process is claimed to provide cement with enhanced durability, as the calcium carbonate precipitation within the CSH gel creates a denser, less porous structure [22]. However, carbonation reduces the alkalinity of the cement paste and in turn can lead to the early corrosion of reinforcing steel, especially in the presence of chloride ions [23]. The combined reliance on high pressure CO₂ infusion and the accompanying acidification of the cement binder with consequent passivation loss limits the application scope of such carbonation methods to pre-cast, unreinforced concrete products, such as bricks, blocks and paving tiles, which together account for no more than 20% of all concrete uses [24]. Such cements, however, could potentially be used with non-steel reinforcement, such as glass fibres.

Unlike carbonation curing, mineralisation through carbon capture serves more than one role; the CO₂ is converted into minerals, such as calcium carbonate and magnesium carbonate, which are used in a multitude of industries. The method of mineralisation can include the carbonation of metal oxide by-products from mining processes [25] or through precipitating the minerals using CO₂ absorption
in an aqueous alkali solution [26]. In the cement industry, calcium carbonate can be used as a clinker replacement, as described in Section 1.1. Based on theory and experience, the maximum efficiency of CO$_2$ mineralisation has been estimated at 68% [27].

The use of mineralised CO$_2$ in the form of PCC-A (produced through carbon CAPCON technology developed by CCM) as a cement admixture is the focus of this paper, where first results have shown that a significant reduction of CO$_2$ emissions of around 25% can be achieved by the substitution of clinker with carbon-negative PCC-A (see Section 3.4). The PCC-A in cement reacts with water during formation to produce hydrated calcium carboaluminates that develop from the hydration and partial carbonation of C$_3$A phases. In doing so, the interaction between water and C$_3$A is modified, leading to a more rapid formation of ettringite. By stabilising carboaluminates, sulfate is forced to remain in the ettringite and so conversion back to monosulfoaluminate becomes thermodynamically unfavourable [8]. This potentially impacts the role of gypsum, a widely used setting retarder [28].

The reference CO$_2$ content of an off-the-shelf PLC was determined in the course of the tests, using XRD. At 11.4%, it has a comparable CO$_2$ content to carbonation cured cements at 11.0% [21], but with none of the inherent disadvantages or limitations. Further testing of PCC-A blended cements is expected to confirm that a significantly higher embodied CO$_2$ content can be achieved without either detriment to quality or limitation to the range of products made and applications deployed, when using these new generation Portland cements.

2. Materials and Methods

2.1. Materials

Two types of cement were used in the tests carried out—a CEM I 52.5N OPC and a CEM II/A-LL 32.5R PLC. Hanson Cement UK supplied both products. The main constituents of the cements were determined through XRD analysis and Rietveld Refinement, as detailed in Sections 2.2 and 3.1. The manufacturer specifies that both cements have a Blaine fineness between 300 and 500 m$^2$/kg.

The PCC-A used is a product of the carbon CAPCON process developed by Carbon Capture Machine (UK) Limited. It was prepared from CO$_2$-rich flue gas, where CO$_2$ was selectively dissolved at the point of emission in dilute aqueous alkali (NaOH) solution. The solution was then mixed with calcium-containing brine to produce the PCC-A, which was subsequently filtered, rinsed and dried before use to remove contaminants, such as NaCl.

2.2. X-ray Diffraction (XRD) Analysis

Samples of the PCC-A, and both cement types, were tested using a PANalytical X’Pert X-ray powder diffractometer to determine the composition. Powder specimens were analysed at room temperature at a 2$\theta$ range from 30° to 80°, Cu K alpha radiation, using a scan period of 10 min.

2.3. Rheological Data

The rheological data of the freshly mixed cement pastes were measured using a continually increasing shear rate up to 200 1/s over a period of 100 s before decreasing at the same rate to 0. A $\varnothing$25 mm coaxial cylinder spindle was used in conjunction with a $\varnothing$40 mm sample pot.

The cement and PCC-A were weighed to provide a total solids content of 200 g and the desired water/solids ratio was also weighed. The PCC-A was added to the water and stirred continuously to ensure it was evenly dispersed. The PCC-A/water slurry was then combined with the cement in a mixing bowl and mixed continuously for 90 s; this mixing period was considered sufficient to ensure the hydration of the cement paste while reducing the risk of early-set formation. Within 30 s of the mixing ending, the paste was transferred to the sample pot and connected to the rheometer and the test was started.

Data were collected for samples of OPC, PLC and PCC-A blended cement ranging from 0 to 20% PCC-A content. As the water/cement ratio decreased, the range of PCC-A cements that were sufficiently workable to test also decreased.
2.4. Compressive Strength

Cement pastes were prepared in the same way as for rheological testing, but with a longer mix time in accordance with BS EN 196-1:2016 [29]. A longer mix time, compared with the rheological tests, was used to ensure the PCC-A was well distributed in the paste before casting. The paste was then cast in 50 × 50 × 50 mm cube steel moulds. Samples were maintained at a constant temperature of 25 °C for 24 h before being demoulded. Samples were thereafter fully submerged in a water bath at an ambient temperature for 7 or 28 days. It was observed during casting that cement pastes containing a high PCC-A content were less workable, i.e., they were more viscous than OPC and at the highest PCC-A content were difficult to cast.

After curing, the samples were wiped dry before being tested. Testing was carried out using a hydraulic ram at a loading rate of 1 mm/min. The applied load was increased until the sample fractured. Data were collected electronically with a measurement of displacement and force recorded 6 times per second. The compressive strength was calculated as a variant of the equation provided in BS EN 196-1:2016 [29] to account for the different cube size tested:

\[ R_c = \frac{F_c}{2500} \]  

where \( R_c \) is the compressive strength (MPa), \( F_c \) is the load recorded at the point of fracture (N) and 2500 is the area (mm\(^2\)) that the load was applied to, i.e., 50 × 50 mm.

2.5. Scanning Electron Microscope (SEM) Imaging

Fragments from the compressive strength testing were collected and used for SEM imaging. Pieces no greater than 10 mm in any dimension were coated using carbon evaporation under vacuum. The device used was a Zeiss GeminiSEM Field Emission Scanning Electron Microscope, which allowed for multiple methods of imaging to be used. Images of the fracture surface were obtained using secondary electron (SE) imaging and electron backscatter diffraction (BSE).

SE imaging is the detection of low energy electrons emitted from the surface of the specimen exposed to the high energy electron beam and can be used to determine specimen topography. BSE is the detection of primary electrons that have been diffracted by an angle greater than 90°. The angle at which the electron has been diffracted is related to the atomic structure of the scattering material and can be used to reveal changes in specimen composition [30].

3. Results and Discussion

3.1. XRD Analysis

XRD diffractograms of the OPC and PCC-A blended cement samples tested were analysed using Rietveld Refinement, using Profex/BGMN [31] in accompaniment with structure data from ICSD [32]. Tables 1 and 2 provide, in standard notation, the phases identified in the cement clinker and the oxides present in those phases. The PLC used for comparison interestingly had a calcium carbonate content of 18.27%, which was more than the maximum amount of PCC-A blended with OPC that was tested (15%).

| Mineralogical Composition | Phase Wt. % | Chemical Composition | Oxide Wt. % |
|--------------------------|-------------|---------------------|------------|
| C\(_{3}\)S                | 59.65       | SiO\(_2\)           | 20.28      |
| C\(_{2}\)S                | 15.24       | Al\(_2\)O\(_3\)     | 4.71       |
| C\(_{3}\)A                | 11.81       | Fe\(_2\)O\(_3\)     | 3.27       |
| C\(_{4}\)AF               | 8.65        | CaO                 | 67.13      |
| C\(_{5}\)S\(_2\)H\(_2\)   | 4.65        | SO\(_3\)            | 2.54       |

MgO 0.67
K\(_2\)O 1.40

Table 1. Quantitative analysis of Ordinary Portland Cement (OPC) used in tests and oxides present.
Table 2. Quantitative analysis of Portland limestone cement (PLC) and oxides present.

| Mineralogical Composition | Phase Wt. % | Chemical Composition | Oxide Wt. % |
|---------------------------|-------------|----------------------|-------------|
| C₃S                       | 60.11       | SiO₂                 | 15.23       |
| C₂S                       | 7.66        | Al₂O₃                | 3.33        |
| C₃A                       | 9.42        | Fe₂O₃                | 2.75        |
| C₄AF                      | 1.03        | CaO                  | 64.40       |
| C₅H₂                      | 3.51        | SO₃                  | 1.41        |
| CaCO₃                     | 18.27       | K₂O                  | 0.83        |
|                           |             | CO₂                  | 11.41       |

The chemical composition of the PCC-A used in the blended cements is summarised in Table 3. This PCC-A was comprised of calcite with a very small, residual salt content remaining from the production process. No other CaCO₃ polymorphs were identified using Rietveld Refinement.

Table 3. Quantitative analysis of PCC-A used in cement blends.

| Compound     | Wt. % |
|--------------|-------|
| Calcite      | 99.8  |
| Aragonite    | 0.0   |
| Vaterite     | 0.0   |
| Halite, NaCl | 0.2   |

The NaCl content attributed to the PCC-A dropped to 0.02% when a 10% blended PCC-A cement was produced. This is the same as the NaCl content in the fresh tap water used to cure the test samples. Although the presence of NaCl can lead to an increased compressive strength, at this very low level it would be limited to no more than ~0.3%. A 10+% relative increase in the 28-day compressive strength of PCC-A blended cements, therefore, cannot be attributed to the presence of NaCl. The logical conclusion is that it is attributed to the use of PCC-A.

XRD analysis provides a powerful tool for identifying the purity of PCC-A samples and will be of great benefit in future when different PCC-A polymorphs and granularities are used.

3.2. Determination of Strength

After 7 days of curing, the PCC-A cement blends showed a reduction in strength with the carbon-negative Precipitated Calcium Carbonate (PCC) content above 4%. However, after 28 days, an increase in strength was observed up to 10% PCC. This increase in strength was accompanied by a decrease in the workability of the cement paste, tentatively attributable to the formation of carboaluminate phases, which act as a strong binder between the cement paste, unreacted PCC-A and, in the case of concrete, aggregates [13].

The complete set of results from the compressive strength testing is presented in Figure 3. It can be seen that an increase in strength is gained after 7 days of curing for PCC-A content between 1 and 4%. Beyond 4% PCC-A, the strength of the cement starts to decrease as the PCC-A content increases. After 28 days, the strength increase resulting from the inclusion of PCC-A continues up to and beyond 15%, reaching a maximum strength of 54.8 MPa at 10% PCC-A content. This represents an increase of approximately 20% compared to the OPC reference. A PLC sample was separately tested and found to have a compressive strength of 18.1 MPa and 22.3 MPa at 7 and 28 days, respectively.
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Since all the tests were carried out on cement pastes, the expected 52.5 MPa compressive strength of the OPC was not reached—this value is normally attributed to mortar strength.

Future work on strength will require testing samples at lower water/cement (w/c) ratios and increasing the curing time up to 365 days. OPC, PLC and 10% PCC-A will also be tested at a similar workability, as is shown in Section 3.3. The use of PCC-A in mortars will be studied, as the formation of carboaluminate phases has been documented to have strong bonding properties between aggregates and cement paste [13].

3.3. Rheological Analysis

Cement pastes with water/cement ratios of 0.5, 0.4 and 0.3 were tested using the method described in Section 2.3. Data were recorded using a Brookfield R/S+ rheometer in conjunction with the Rheo3000 software package. The addition of PCC-A to cement created a more viscous paste (Figures 4–7), whereas the PLC tested for comparison was less viscous (Figures 4, 6 and 7).
The overt rise in viscosity is termed “false set” and can sometimes be overcome by using admixed plasticiser. The false-setting compositions, assuming that normal strength gain occurs at later stages, paste starts early. However, the pastes then break down quickly and become fluid and workable again. Figure 3), cement pastes of OPC and PLC that provided a similar workability by adjusting the w/c however, these compositions would not be useful in conventional construction. They might actually be useful in developing sprayed concretes—exploring this is a task for the future.

As the w/c ratio of the cement pastes decreases, the PCC-A cement samples over 10% became too viscous for the rheometer to start and are not included in Figures 6 and 7. This is likely due to the large shear stress that can be observed in the initial part of the tests, where setting of the cement paste starts early. However, the pastes then break down quickly and become fluid and workable again. The overt rise in viscosity is termed “false set” and can sometimes be overcome by using admixed plasticiser. The false-setting compositions, assuming that normal strength gain occurs at later stages, might actually be useful in developing sprayed concretes—exploring this is a task for the future. However, these compositions would not be useful in conventional construction.

As the 10% PCC-A cement samples showed the greatest compressive strength of those tested (see Figure 3), cement pastes of OPC and PLC that provided a similar workability by adjusting the w/c...
ratio were found. These w/c ratios are shown in Figure 8. The volume of water required for a 10% PCC-A to obtain a similar workability to OPC is 25% higher. For PLC, the required volume is 2.5% lower. Further work is clearly needed to support the wide use of PCC-A cement blends in future.

Figure 7. Shear stress in cement samples containing w/c = 0.3. The 10% PCC-A samples exhibit similar rheological properties.

Figure 8. Workability of three different blends to determine the w/c ratio for which OPC, PLC and 10% PCC-A samples exhibit similar rheological properties.

Further work is planned and will be carried out on the rheology of PCC-A cement blends to definitively establish why they behave in a manner dissimilar to what is known for PLC.

3.4. Scope for Emission Reduction of PCC-A Cement Blends

The reduction in CO₂ emissions through the use of PCC-A can be calculated as follows:

\[
\text{CO}_2 \text{ reduction, } C = w_{\text{opc}} \cdot m_{\text{opc}} + w_{\text{pcc}} \cdot m_{\text{pcc}},
\]

where \(w_{\text{opc}}\) and \(w_{\text{pcc}}\) are the mass fractions of OPC and PCC-A respectively, and \(m_{\text{opc}}\) and \(m_{\text{pcc}}\) is the mass, in grams, of the CO₂ emitted in the production of 1 kg of the respective product. The final potential reduction in cement is then calculated using the reduction in clinker required to achieve the same compressive strength as OPC. The final reduction in CO₂ emissions is then found as:

\[
\% \text{ reduction } \text{CO}_2 = (1 - (C_i \cdot (R_{\text{c-opc}}/R_{\text{c-i}}))/C_{\text{opc}})) \cdot 100,
\]

where \(C_i\) is the CO₂ reduction of a PCC-A with \(i = 1–15\) wt.% PCC-A, \(C_{\text{opc}}\) is the CO₂ emitted from OPC production, \(R_{\text{c-opc}}\) is the mean compressive strength of OPC and \(R_{\text{c-i}}\) is the mean compressive strength of the PCC-A blended cement with \(i = 1–15\%\). For example, the reduction in CO₂ for the 10% PCC-A cement is calculated as follows:

\[
\text{CO}_2 \text{ reduction, } C_{10\%} = 0.9 \cdot 951.5 \text{ gCO}_2/\text{kg} + 0.1 \cdot (-100) \text{ gCO}_2/\text{kg},
\]

\[
C_{10\%} = 846.35 \text{ gCO}_2/\text{kg},
\]

\[
\% \text{ reduction } \text{CO}_2 = (1 - (846.35 \text{ gCO}_2/\text{kg} \cdot (44.8 \text{ MPa}/54.8 \text{ MPa}))/951.5 \text{ gCO}_2/\text{kg})) \cdot 100,
\]

\[
% \text{ reduction } \text{CO}_2 = 27.3\%.
\]
Traditionally, the production of one kilogram of OPC clinker emits on average ~950 gCO₂ [33]. The production of one kilogram of PCC-A sequesters ~100–350 gCO₂, depending on the alkali used in the capture process. To date, the carbon mitigation of two alkalis has been examined—sodium hydroxide and, to a limited extent, ammonia. The production of sodium hydroxide has a carbon intensity (i.e., the weight ratio of CO₂ emitted in production per unit weight of product made) of ~870 gCO₂/kg [34] and ammonia has a carbon intensity of ~1,600 gCO₂/kg [35]. However, there is a body of evidence to suggest that 80% of ammonia used in the reaction could be regenerated and reused, and so the carbon intensity could be reduced to 667 gCO₂/kg due to its lower material demand during the CO₂ capture process. For further information, see the Technology Centre Mongstad (TCM) Chilled Ammonia Process (CAP), which is based on the chemistry of the NH₃-CO₂- NH₃ system [36].

The final CO₂ sequestered using either sodium hydroxide or ammonia is then 100 gCO₂ or 350 gCO₂, respectively. The PCC-A used in the test results that have been reported was produced using sodium hydroxide, and the values in Figure 9 reflect that. Figure 10 shows the potential reduction in emissions if recovered ammonia is used as the alkali in the PCC-A production process. Alkalis from waste could potentially offer equally low carbon intensity alternatives.

![Figure 9](image1.png)  
**Figure 9.** Clinker substitution with PCC-A produced using NaOH: (a) CO₂ emissions of PCC-A cement blends with increasing PCC-A content and strength considerations, and (b) percentage reduction in CO₂ emissions of PCC-A cement blends through clinker substitution and strength gain.

![Figure 10](image2.png)  
**Figure 10.** Clinker substitution with PCC-A produced using NH₃: (a) CO₂ emissions of PCC-A cement blends with increasing PCC-A content and strength considerations, and (b) percentage reduction in CO₂ emissions of PCC-A cement blends through clinker substitution and increase in strength.

The introduction of PCC-A blended cements could have a significant impact on the carbon footprint of the cement industry (and with that the built environment), due to the reduction in CO₂
emissions in the PCC-A production process and the subsequent substitution of clinker using PCC-A. As shown in Figure 9, the 15% PCC-A blended cement made using a NaOH CAPCON process provided a reduction in emissions of 27% compared to OPC. A PCC-A made using an ammonia process would deliver a 30% reduction. Future work on the use of PCC-A will involve a detailed cradle-to-grave Life Cycle Assessment (LCA) of PCC-A cement blends, and it will study how they compare against other cements and cement blends in terms of overall performance, resistance to deterioration and cost.

3.5. SEM Imaging

SEM micrographs were obtained using a secondary electron detector. Figure 11 is of the non-optimised PCC-A that was used to produce the samples and test results reported in this paper, at progressively higher magnifications. Figure 12 presents and the backscatter electron detector images of the fracture surface of the 15% PCC-A cement blend. Both large and small PCC-A crystals were identified on the fracture surface of the sample using SEM imaging, the latter possibly due to the increased surface area for reaction required for the formation of carboaluminate phases. Figure 13 is a backscatter detector image of the fracture surface of the 10% and 15% PCC-A cement blends.

![SEM micrographs of the PCC-A that was used in cement blends using secondary electron (SE) detection. Grain size was estimated to range from less than 100 nm to 20 µm. (a) 200× magnification of PCC-A and (b) 4000× magnification of the same PCC-A sample.](image1)

![SE images showing fracture surface topography in a sample of the 15% PCC-A cement blend at different magnifications. (a) 4000× magnification and (b) 10,000× magnification.](image2)
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Progression of phase composition is planned for samples cured from 1, 7, 28 and 365 days.

Conflicts of Interest: The authors declare no conflict of interest.

Electron Microscopy was performed in the ACEMAC Facility at the University of Aberdeen.

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4. Conclusions

PCC-A, made using flue gas CO₂ in CCM’s CAPCON process, used as a Portland cement admixture, leads to an increased compressive strength that peaks at 10 wt.% PCC-A content. Combined with the carbon-negative nature of PCC-A, the resulting reduction in the clinker factor and the cumulative carbon content, the carbon footprint of a PCC-A blended cement containing 15% PCC-A is equivalent to ~73% of that of the OPC average.

The strengths of PCC-A cement blends increase with PCC-A content, but the workability reduces, and, speculation notwithstanding, a full and detailed understanding of the processes that are involved is needed.

To achieve the same level of workability as an OPC paste, a PCC-A cement paste requires 25% more water, which is not how a PLC cement behaves (requiring 2–5% less water).

Future work to be undertaken on the effects of PCC-A in blended cements will include:

- The testing of PCC-A for use in mortars to determine the binding effects on aggregates;
- Examination of PCC-A cements after long curing times to study the evolution of carboaluminate phases;
- Investigation of the inverse effects on rheology compared with PLC;
- Full LCA of PCC-A blended cements using more alkali-efficient mineralisation processes;
- Preparation of PCC-A blended cements containing polymorphs of PCC-A to determine if crystal structure plays a role in reactivity during hydration.

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