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4 Supercritical carbonation of calcareous composites: Influence of curing
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26 Abstract
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28 This paper reports the effect of curing on the susceptibility of cementitious composites to
29 carbonation using supercritical carbon dioxide. Samples made using a compression
30 moulding technique were cured in water before and/or after carbonation and the effect on
31 porosity, microstructure, solid phase assemblage and flexural strength was determined. In
32 terms of development of mechanical strength, no benefit was gained from any period of
33 pre- or post-carbonation curing regime. Yet samples cured prior to carbonation
34 underwent minimal chemical reaction between supercritical carbon dioxide and calcium
35 hydroxide, unhydrated cement or C-S-H. Thus there was no correlation between chemical
36 degree of reaction and strength development. The effects responsible for the marked
37 strength gain in supercritically carbonated samples must involve subtle changes in the
38 microstructure of the C-S-H gel, not simple pore filling by calcium carbonate as is often
39 postulated.
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51 Keywords
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53 Super-critical carbonation; mechanical processing; lime; cement; flexural testing;
54 petrography; curing
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4 1. Introduction
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8 The effect of scCO₂ on cementitious materials was first investigated by the oil industry in
9 the context of deep well linings, but its use as a treatment to improve the properties of
10 cement composites was initiated by Jones in late 1990s [1-3]. Supercritical carbonation
11 (SCC) of cementitious composites is chemically similar to natural carbonation in that the
12 carbon dioxide diffuses into the capillary pores of the cement paste and combines with
13 water present in capillary pores, forming carbonic acid (H₂CO₃). This acid dissociates
14 into carbonate (CO₃⁻²) and hydrogen (H⁺) ions which then react with the portlandite
15 (Ca(OH)₂) formed during hydration, cement hydrates, any unhydrated cement (mainly
16 C₃S and C₂S), and residual sodium and potassium ions in the pore solution of the cement
17 matrix. In calcareous composites, the binder is formed from a mixture of cement and/or
18 other calcium-bearing additive materials such as slaked lime, fly ash, steel slag etc. The
19 latter of these also contain a certain amount of active silica. These are generally added to
20 improve technical properties, or to reduce environmental impact and/or cost by partially
21 replacing cement binder with cheaper, potentially ‘low energy’ materials. In in most
22 cases, sufficient siliceous material is retained in the matrix such that the principal binding
23 phase remains C-S-H gel (in contrast to pure lime mortars). The scCO₂ will also react
24 with the calcium-bearing phases in these additive materials.
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41 These reactions form calcium carbonate, hydrated silica (as a result of decalcification of
42 C-S-H gel) and other minor products including sodium and potassium carbonates and
43 hydrated alumina [4-8]. Compared to natural or ordinarily accelerated carbonation
44 however, SCC is greatly accelerated, with complete carbonation of engineering-sized
45 components being achieved in hours rather than years. This is mainly attributable to the
46 significant rise of CO₂ solubility in the pore solution, and the relative ease with which
47 high-density scCO₂ can penetrate and diffuse into the cement paste pore network [7,8].
48 As with natural carbonation, the process reduces the hydroxide concentrations in the pore
49 solutions, lowering its alkalinity from pH = 12.5-13.5 in uncarbonated cement paste
50 down to pH ≈9 in the fully carbonated zone. The exact pH will depend on the HCO₃/CO₃
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4 equilibria, which in turn will be controlled by both the degree of carbonation and amount
5 of alkalis (NaOH, KOH) present [4,9,10].
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10 The carbonation process also alters the microstructure of calcareous based composites,
11 [1,7,8,11-15]. The molecular volume of calcium carbonate (normally calcite) is 11.2%
12 greater than that of calcium hydroxide. Thus the calcite precipitated within the hardened
13 cement paste (hcp) matrix during carbonation (both indirectly from the decalcification of
14 C-S-H gel and accelerated hydration/carbonation of residual unhydrated cement particles,
15 and directly from carbonation of the free calcium hydroxide dispersed through the hcp)
16 fills up pores with small tightly packed crystals of calcite. This reduces the total pore
17 volume, pore size and permeability of the composite [4,6,10] and also enhances the
18 compressive and flexural strength [7,8,10,16,17]. Accordingly, much recent work has
19 focussed on documenting the improvements in the microstructural, mechanical and
20 durability properties of cementitious composites afforded by treatment with scCO₂ [6,10-
21 15,18-20]. Some researchers have investigated combination of novel production methods
22 combined with SCC with a view to enabling mass-production of high-value functional
23 and/or structural SCC ceramic composites from a range of starting materials including
24 cement, lime, various aggregates and waste materials (e.g. PFA, steel slag) [21,22].
25 Aggregate chemistry and grading, binder composition and aggregate-binder ratios were
26 all found to have significant effects on the properties of SCC cementitious composites.
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42 The work reported in this paper was focused on understanding the effect of conventional
43 wet curing of (as commonly used for pre-cast concrete components) on: the rate and
44 extent of carbonation; the flexural strength; and the microstructure of supercritically
45 carbonated specimens of cementitious composites. For the SCC-treated samples, the
46 effect of two curing scenarios was examined; curing prior to and curing after SCC
47 treatment. This was done to examine if any technical benefit could be gained from such
48 curing either before or after specimens are exposed to scCO₂. These specimens were
49 compared with uncarbonated, conventionally cured samples analogous to those currently
50 manufactured in industry. The effect of these combinations of SCC treatment and moist
51 curing was investigated using X-ray diffraction (XRD), combined differential thermal
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4 analysis with thermo-gravimetric analysis (DTA/TG), thin section petrography (TSP),
5 helium pycnometry and a 4-point flexural strength test method.
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10 2. Experimental methods 11 12

13 2.1. Materials 14 15 16 17

18 All specimens were manufactured with an aggregate to binder ratio of 5:2. The binder
19 was 1.5:0.5 ordinary Portland cement [C] : hydrated lime [L] by mass; aggregate [A] was
20 65:35 by mass of crushed limestone : silica sand. These combinations were found to be
21 optimal in previous work [23] with regard to ease of mechanical processing, reliability of
22 carbonation and development of strength. The lime (Buxton Lime, UK) used comprised
23 of >96% Ca(OH)_2 as determined using X-ray fluorescence (XRF) and the remainder
24 made up of CaCO_3 , CaO , SiO_2 and Mg(OH)_2 . The Portland cement (Tarmac, UK)
25 comprised 65% CaO , 20.25% Si_2O , 5.3% Al_2O_3 , 3.25% Fe_2O_3 and 3.3% SO_3 , as
26 determined by XRF and the remainder made up of MgO and K_2O . The crushed limestone
27 [CL] (Buxton Lime/Tarmac, UK) comprised 98% CaCO_3 with upper:lower 20 percentile
28 particle size of 2.0:0.15 mm and pure silica sand [S] (Buxton Lime/Tarmac, UK) had
29 upper:lower 20 percentile particle size of 0.5:0.15 mm).
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42 2.2. Sample Preparation 43 44 45

46 Green samples (i.e. fresh, uncured and uncarbonated specimens with only sufficient
47 strength for handling purposes) were manufactured using a compression moulding
48 technique combined with vacuum dewatering. A two-part perforated stainless steel tool
49 (Figure 1) was used to produce six trapezoidal specimens per sample (length: 170 mm,
50 width: 22/34 mm, depth: 17 mm) in each operation, using 9 MPa of pressure for 1 min,
51 preliminary work having established this as the optimum processing conditions [23].
52 Excess water squeezed out of the samples was removed with a vacuum pump connected
53 via a manifold to the perforations in the tool. The tool was lined with filter paper to
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4 prevent egress of solid material. Surplus samples were used to estimate the quantity of
5 water removed during pressing. The water to binder ratio (w/c) of the cementitious paste
6 within the 1.5C:0.5L:5A composite for the compression moulding process was 0.55.
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8 After pressing the w/c of manufactured green specimens was reduced to 0.3.
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13 [Figure 1]
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15 16 17 2.3. Wet curing and pre-conditioning and supercritical carbonation. 18 19

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21 Three distinct post-pressing treatment phases were used:

- 22 • Pre-conditioning: a partial drying treatment design to optimise moisture levels in
23 the green specimens to promote carbonation [12,13,24]. Samples were dried in a
24 fan oven for 12 h at 25°C to remove ~75% of the free water.
- 25 • Curing: immersion in water at room temperature for 3, 7 or 28 days.
- 26 • SCC treatment: exposure to static water saturated scCO₂ at 60°C, 10 MPa for 24 h
27 in a sealed stainless steel pressure vessel (see Farahi et al [22] for further details
28 of the experimental set up). Corresponding control samples were stored sealed at
29 room temperature during the 24 h period in which treated samples were
30 undergoing SCC treatment.
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41 These were combined in 7 sequences of experiments: 1 basic (C1), 2 pre-cured (C3, C7),
42 1 double-cured (C7-28) and 3 post-cured (C3*, C7*, C28*). Each sequence comprised a
43 treated set exposed to SCC treatment, and a control set stored sealed at room temperature
44 for 24 h. These are described below:
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- 48 • C1: Preconditioning followed by either SCC treatment (treated) or stored sealed
49 (control);
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- 52 • C3: Curing for 3 days then preconditioning, followed by either SCC treatment
53 (treated) or stored sealed (control);
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- 56 • C7: as C3 but with 7 days of curing;
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- 58 • C7-28: as C7 but followed by a further period of post-treatment curing for 28
59 days;
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- C3*: as C1 but followed by post-treatment curing for 3 days;
- C7*: as C1 but followed by post-treatment curing for 7 days;
- C28*: as C1 but followed by post-treatment curing for 28 days.

At the end of each combination, samples were tested for mechanical and microstructural properties.

2.4. Mechanical Testing

Triplicate specimens were tested for flexural strength using a fully articulated 4-point bending fixture attached to a screw controlled machine (Testometric Micro 100KN PCX). All testing was performed using a small load cell (100 kgf) and at a constant cross head displacement of 1 mm min^{-1} . Mid span deflection was measured using an integrated LVDT-type transducer. The machine was controlled by computer software which captured all load – displacement data with 0.1 N and $1 \text{ }\mu\text{m}$ resolution up to failure. Flexural strength at failure (modulus of rupture) was calculated using standard beam theory.

2.5. Chemical Analysis

2.5.1. X-ray diffraction (XRD)

Immediately after mechanical testing, the degree of carbonation of treated specimens was analysed using XRD (Philips PW 1830). Small portions from each specimen were taken and finely ground to pass a $150 \text{ }\mu\text{m}$ sieve. The analysis was then performed using $\text{CuK}\alpha$ radiation between 15° and $80^\circ 2\theta$ at $0.6^\circ \text{ min}^{-1}$ (0.02° per step, 2 s per step). For each scenario, triplicate control and carbonated specimens were tested and degree of carbonation was examined semi-quantitatively by computing the mean ratio of peak height of two most prominent peaks for portlandite (CH) and alite (C_3S) for the three control and three carbonated samples.

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8 Combined differential thermal analysis/ thermo-gravimetry (DTA/TG, PL-STA 1500),
9 was used as a quantitative technique to determine the nature and relative concentrations
10 of various compounds present in the samples such as calcium carbonate (CĈ) and
11 calcium hydroxide (CH). DTA-TG can also provide some information on the nature and
12 relative quantity of compounds such as C-S-H gel that are amorphous and thus could not
13 be recognised by XRD [25]. For this test, specimens were ground to pass a 150 µm sieve
14 and then immediately analysed in air, between 20 to 1100°C at a heating rate of 20°C
15 min⁻¹.
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26 2.6. Helium Pycnometry
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30 Helium Pycnometry was used to investigate the influence of the curing regimes and
31 supercritical carbonation on the pore structure. For each case, 10×13×20mm cuboids
32 were cut from mechanical testing remnants using a precision saw and dried by immersion
33 in acetone for 48hr, followed by storage over silica gel until reaching constant weight
34 (taking between 14 and 21 days) after the method described by Aligizaki [26]. The
35 cuboids were then accurately measured by micrometer and weighed, providing the bulk
36 density. They were then tested in a helium pycnometer (AccuPyc 1330, Micromeritics)
37 over 10 purge cycles to provide the true densities, from which the porosity was
38 calculated. Samples were tested in triplicate.
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48 2.6. Thin section petrography
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52 Thin section petrography (TSP) was used to investigate the effect of mix design, curing
53 and SCC on microstructure. TSP is a more useful method for examining the
54 microstructure of a composite than SEM when authoritative identification of phases (eg
55 differentiation between CH and CĈ) is required [12,26]. Standard 30 µm thick, clear
56 resin impregnated thin sections were prepared. The slides were then examined under a
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4 modified Vickers petrological microscope, using plane polarised, dark field plane
5 polarised and/or crossed polarised light.
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10 11 3. Results 12

13 14 15 3.1. Bending strength evaluation 16

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19 Figure 2 shows the variation in flexural strength of control vs scCO₂ treated samples with
20 curing conditions. As expected, SCC treatment results in a significant improvement in
21 strength over control samples in all cases. However, no significant further benefit could
22 be gained from any period of water curing prior to carbonation; only slight strength
23 improvements were observed in specimens cured after carbonation. Thus, from a
24 practical point of view, carbonation immediately after demoulding and pre-conditioning
25 would be the preferred production option. The flexural strength of samples thus treated (C1,
26 13.7 MPa) was about 60% higher than that achieved by normal 28 days conventional
27 curing (C28* control, 8.7 MPa).
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37 [Figure 2]
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44 XRD results (Figure 3, 4) showed complete depletion of crystalline CH and C₃S in
45 samples carbonated immediately after pre-conditioning (C1, C3*, C7*, C28*), suggesting
46 that treated samples were fully carbonated. For treated samples only wet-cured after
47 carbonation (C3*, C7*, C28*), CH did not reappear in samples after they were exposed
48 to water, suggesting that no amorphous C₃S (which would not be detected by XRD) is
49 left after SCC treatment that can enable secondary hydration.
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57 However, treated samples wet-cured before SCC treatment (C3, C7, C7-28) were
58 resistant to carbonation and failed to become fully carbonated; significant amounts of CH
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4 and C₃S were detected in the samples after SCC treatment. More CH and C₃S remained
5 in treated samples (compared to relevant control sample) exposed to longer curing prior
6 to carbonation; in samples cured for 3 and 7 days prior to carbonation (C3, C7) CH and
7 C₃S were depleted by 53% or 35 – 40% respectively by SCC treatment.
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13 In samples exposed to both pre- and post-treatment curing (C7-28), the C₃S remaining
14 after SCC treatment appeared to have its ability to further hydrate restricted, since the
15 proportion of remaining C₃S further hydrated during post-treatment curing was much
16 smaller for carbonated samples cf. control samples. Interestingly, the strength of treated
17 C7-28 samples was the highest recorded (14.9 MPa) yet the degree of carbonation (as
18 measured by CH and/or C₃S depletion) was lowest. These results confirm that traditional
19 measurements of degree of carbonation (e.g. phenolphthalein indicator tests) are unlikely
20 to be a useful guide to strength gains – and hence microstructural development – in
21 carbonated samples; the rarely measured effect of carbonation on the gel phases (i.e. C-S-
22 H) is likely to be much more important.
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33 The amount of C₃S present in control samples was reduced as curing times increased, as
34 expected.
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39 [Figure 3]

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41 [Figure 4]

42 43 44 3.3. DTA/TG

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47 DTA/TG was used primarily to investigate samples cured prior to carbonation, in order to
48 investigate the source of strength enhancement despite incomplete carbonation of
49 crystalline phases. Figure 5 shows the DTA thermograms for C1, C3, C7-28 and C28*.
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55 The endotherms between 100-200°C reflect the dehydration of ‘low temperature
56 hydrates’ (LTH) i.e. C-S-H gel and AF_t phases. For control samples, the area and
57 complexity of the LTH peak increases with curing time as expected, reflecting continued
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4 hydration of the cement. The effect of SCC treatment on the LTH peak is markedly
5 different depending on the curing sequence. For samples not exposed to pre-treatment
6 curing, SCC treatment (C1, C28*) almost removes the LTH peak, suggesting that the gel
7 phases are significantly affected (probably by decalcification and dehydration to form a
8 silica gel). In contrast, for samples cured before treatment (C3, C7-28), the effect of SCC
9 treatment on the LTH peak is much less marked and lessens with increased curing time;
10 for C3, some broadening of the LTH peak can be observed but for C7-28 the peak
11 remains essentially unchanged, retaining its sharpness and shoulders.
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21 The endotherm between 450-530 °C is due to the dehydroxylation of CH. In the cases
22 where specimens were carbonated immediately after manufacturing (C1, C28*) CH has
23 been depleted fully by the treatment. In the treated samples exposed to curing prior to
24 carbonation (C3 and especially C7-28) a considerable amount of CH remained in the
25 samples after SCC treatment; the longer they remained in water (C7-28 cf. C3), the more
26 resistant they have become to carbonation of CH. These results confirm the XRD studies
27 (Figure 3).
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35 The small blip at ~ 580 °C is caused by the $\alpha \rightarrow \beta$ crystal form inversion in the quartz sand
36 aggregate [12].
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41 The effects in the DTA thermograms at 650 °C onward are related to decomposition of
42 calcium carbonate ($\text{C}\hat{\text{C}}$). The contribution to these effects from the aggregate (65%
43 limestone) tends to obscure those associated with development of $\text{C}\hat{\text{C}}$ from carbonation
44 of CH, C_3S and C-S-H and thus no firm conclusion can be drawn. In this regard;
45 quantitative TG analysis is required and this is shown in Table 1 (details of the
46 calculations are given as a footnote to the table). The analysis confirmed a significant
47 increase in $\text{C}\hat{\text{C}}$ content in samples not cured before SCC treatment (C1, C28*), but little
48 or no calcite development in those cured before SCC treatment (C2, C7-28). TG analysis
49 also confirmed that a significant amount of CH remained in samples cured before SCC
50 treatment (C3, C7-28) while samples exposed immediately to scCO_2 (C1 and C28*), CH
51 was shown to be removed completely, confirming the results of the XRD study. For the
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C3 samples, the degree of CH depletion according to XRD (~50%) was somewhat less than that measured using TG (~70%), suggesting that amorphous CH present in the hcp carbonated preferentially. As with the XRD analysis, strength development was not necessarily correlated with degree of carbonation.

[Figure 5]

[Table 1]

Table 1: Quantitative analysis of TG traces for treated and control samples exposed to following curing regimes. Samples made from; 1.5C:0.5L:5A (A=65:56 CL:S)

'Wet' Curing regimes	LTH (%)		Ca(OH) (% w/w)		CaCO ₃ (% w/w)	
	Treated	Control	Treated	Control	Treated	Control
C1	1	1.59	0	4.73	57.59	47.4
C3	2.44	2.39	1.64	5.43	50.98	48.76
C7-28	3.63	3.52	5.75	6.99	45.04	45.04
C28*	1.29	3.41	0	6.99	57.84	47.15

LTH (%): Mass loss (%) recorded between 20 and 200°C. Note: no correction for background drift.
Ca(OH)₂ (%): CH content determined from the mass loss on TG traces between 450 and 530 °C
CaCO₃ (%): Calcite content determined by the mass loss on TG traces between 650 and 875 °C.

3.4. Porosity measurement

Helium pycnometry was carried out the same sample subset as DTA/TG (C1, C3, C7-28 and C28*). Results (Figure 6) suggested that in all cases, the total porosity of the composite was reduced to 0.08 ± 0.01 by the SCC treatment, which would partially account for the increase in strength observed. However, for samples with well-developed pre-treatment microstructure (i.e. C7-28), relatively small changes in porosity (-8%) lead to large increases in strength (+34%); also, where untreated and SCC treated samples have very similar porosity (C28* control and/or C7-28 control, cf. C3 treated), the SCC treated sample has much higher strength (9 – 10 MPa, cf. 14.2 MPa respectively, see Figure 2). This suggests that the strength enhancement effect of SCC treatment is more profound than a simple reduction in porosity; the intrinsic strength of the matrix is also significantly increased.

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6 Increased degree of water curing decreased the porosity of control samples, as expected.
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10 [Figure 6]
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13 3.5. Petrography 14 15

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17 Figure 7 shows the microstructure of an uncarbonated, minimally cured (C1 control)
18 sample. Typical limestone (L) and sand (i.e. quartz, Q) particles have been marked. The
19 groundmass – the material between distinct particles – is generally dark, owing to the
20 poorly crystalline residual unhydrated cement minerals and some CSH gel, studded with
21 bright flecks of small calcium hydroxide crystals and the occasional larger CH crystal
22 (marked CH). The interface between the aggregate particles and the groundmass is not
23 intimate and some porosity can be observed. Since the control sample is uncured and
24 unhydrated, the space at aggregate/matrix interface is largely free of hydration products.
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28 In carbonated samples (Figure 8, C1 treated) however, the groundmass is much lighter
29 and featureless, composed of cryptocrystalline calcium carbonate (identifiable by its high
30 birefringence) mixed with an amorphous phase, presumably decalcified CSH gel. The
31 dark inclusions are pseudomorphs of unhydrated cement grains, which have been
32 carbonated. Compared to control samples, the porosity within the groundmass has
33 reduced considerably, correlating with the helium pycnometry results. There is also very
34 little porosity at the interface between aggregate particles and groundmass; the interface
35 between the limestone and the carbonated groundmass (CG) is now quite indistinct in
36 places.
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50 [Figure 7]
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52 [Figure 8]
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55 Figure 9 shows the microstructure of a C7-28 SCC-treated sample (note the change of
56 scale); the section was taken to include sample surface exposed to scCO₂, which can be
57 seen at the bottom of the figure. The penetration and reaction of the scCO₂ has clearly
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4 been impeded by the pre-cured microstructure; only a relatively shallow (~0.5 mm) layer
5 near the surface has fully carbonated groundmass (CG), while deeper in the sample it
6 remains largely uncarbonated (UG).
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11 [Figure 9]
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15 4. Discussion and conclusions. 16 17

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19 The results from the flexural testing clearly demonstrate that from the point of view of
20 developing mechanical strength, there is no benefit in any period of pre- and/or post-
21 treatment curing. SCC treatment immediately after pressing and pre-conditioning imbues
22 strength well above that which could be expected to be developed by curing alone, and
23 does so within hours rather than days.
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30 Nonetheless, the pre-curing experiments have uncovered an interesting phenomenon;
31 namely that there is almost no correlation between the ‘degree of carbonation’ – i.e. the
32 extent to which CH, residual unhydrated cement and C-S-H gel have reacted with CO₂ –
33 and the strength of the samples. For example, in the C7 and/or C7-28 samples (compared
34 to those remaining uncured prior to SCC treatment), XRD and TG indicated that CH and
35 residual C₃S were minimally depleted by SCC; DTA/TG indicated that the effect of SCC
36 on the gel phase was minimal; pycnometry recorded only a small drop in porosity owing
37 to SCC treatment; and petrographic images clearly show that the so-called ‘carbonation
38 front’ has penetrated only ~0.5 mm into the sample. Yet the strength developed in these
39 apparently minimally carbonated samples was similar to that developed in fully-
40 carbonated samples.
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52 The strength-bearing phase in a cement composite matrix is the C-S-H gel, whether in a
53 ‘pure’ cement-binder system, one modified with lime (as studied here), and/or modified
54 with siliceous additions; thus it is understandable that depletion of crystalline phases is
55 not coupled to strength increases. However, DTA and petrography of minimally-
56 carbonated samples both suggest that the C-S-H is not significantly altered chemically
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4 (i.e., fully decalcified) or morphologically by SCC treatment; or, put another way, the
5 subtle changes in C-S-H microstructure responsible for the strength increase are not
6 detectable by these methods. Previous work by Short et al. [15] using NMR suggested
7 that SCC treatment can fundamentally change the nature of the silica polymerisation in
8 C-S-H gel and this may well be the effect that is at work here. The changes are clearly
9 fundamental (i.e. associated with intrinsic physico-chemical changes in the matrix), as the
10 strength is increased independently of the porosity. This is interesting since many
11 literatures [3-8,18,19] explain the enhancement in strength of carbonated cementitious
12 matrices purely by reduction of porosity through formation of $\text{C}\hat{\text{C}}$ crystals and their
13 precipitation in the matrix pores; clearly this is not the case in this instance. Furthermore,
14 the C-S-H phase must be much more susceptible to SCC carbonation than the crystalline
15 phases. The very thin carbonated layer seen in Figure 9 is insufficient to account for the
16 increase in strength in these samples, implying that in fact scCO_2 has diffused much more
17 deeply into the sample and partially reacted with the C-S-H in the bulk of the matrix
18 relatively quickly. The surface reaction with CH (and, to a lesser extent, unhydrated
19 cement) has proceeded much more slowly and indeed may have been self-limiting via the
20 formation of a protective layer. Clearly, the traditional notion that a carbonation front
21 proceeds in an orderly fashion through a cement-based calcareous composite sample
22 during SCC is deficient. This may also have implications for the study of environmental
23 carbonation.
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Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing – Table 1

‘Wet’ Curing regimes	LTH (%)		Ca(OH) (% w/w)		CaCO ₃ (% w/w)	
	Treated	Control	Treated	Control	Treated	Control
C1	1	1.59	0	4.73	57.59	47.4
C3	2.44	2.39	1.64	5.43	50.98	48.76
C7-28	3.63	3.52	5.75	6.99	45.04	45.04
C28*	1.29	3.41	0	6.99	57.84	47.15

LTH (%): Mass loss (%) recorded between 20 and 200°C. Note: no correction for background drift.
Ca(OH)₂ (%): CH content determined from the mass loss on TG traces between 450 and 530 °C
CaCO₃ (%): Calcite content determined by the mass loss on TG traces between 650 and 875 °C.

Figure 1

Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing – Figure 1

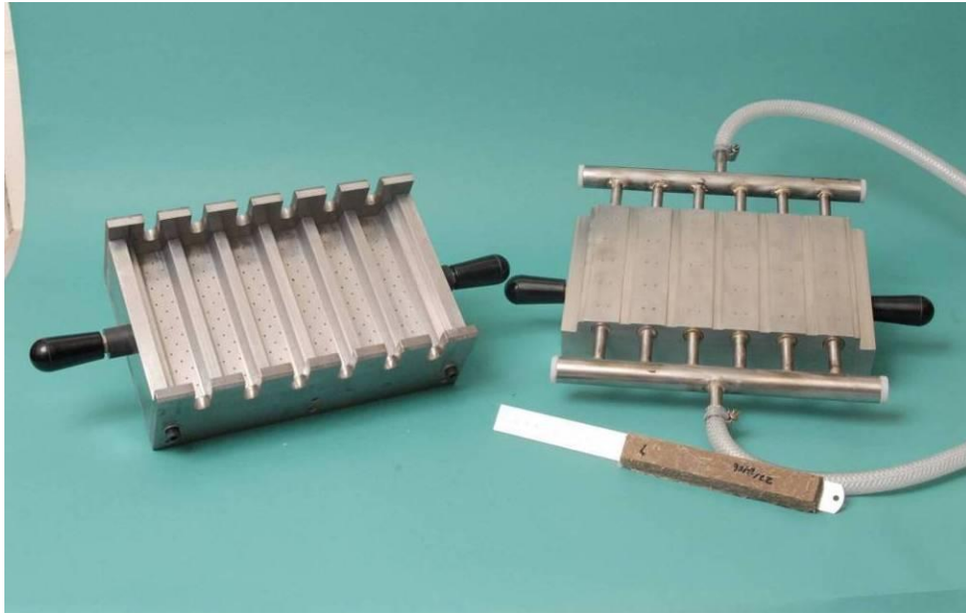


Figure 2

Farahi & Purnell 2012 **Supercritical carbonation of calcareous composite: Influence of curing** – Figure 2.

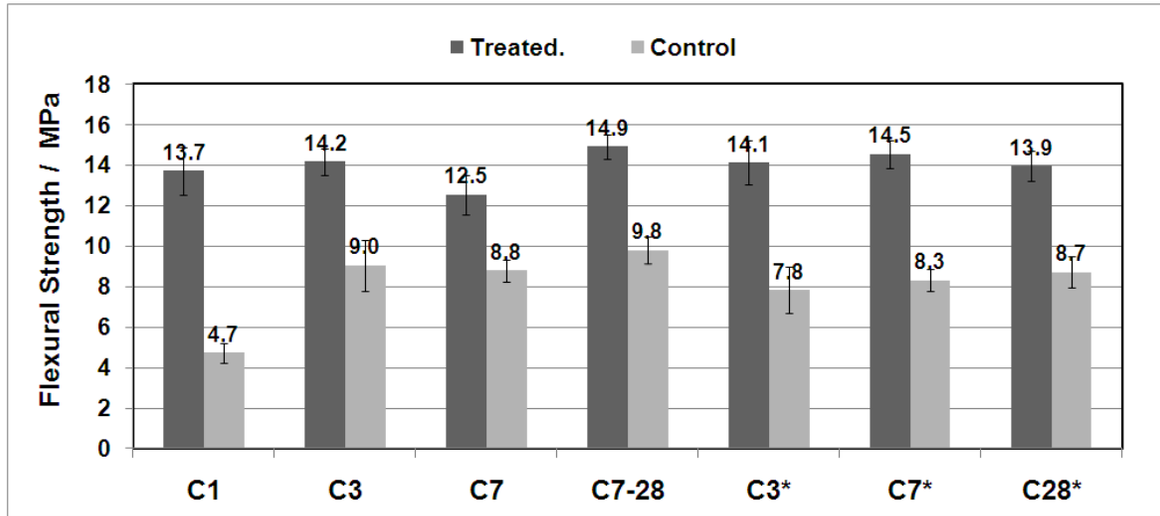


Figure 3

Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing - Figure 3

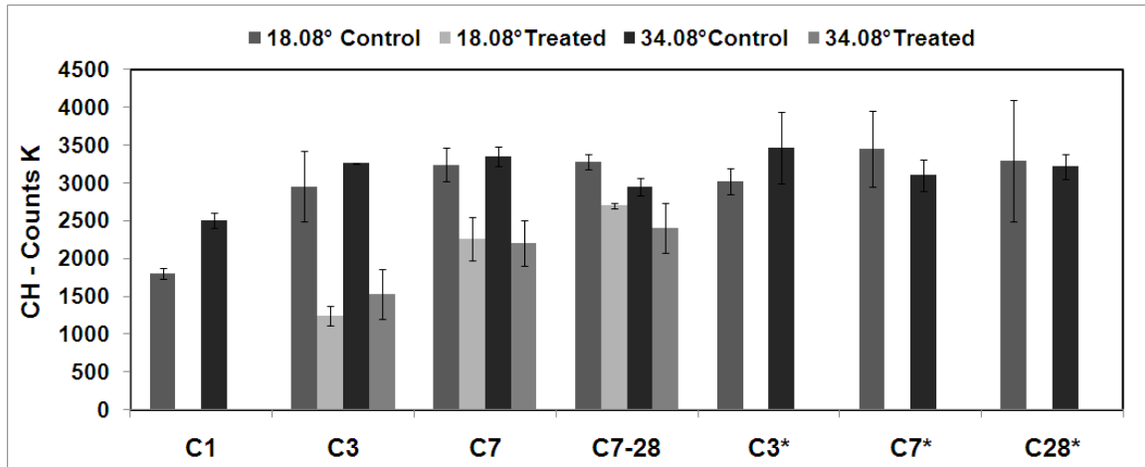


Figure 4

Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing – Figure 4

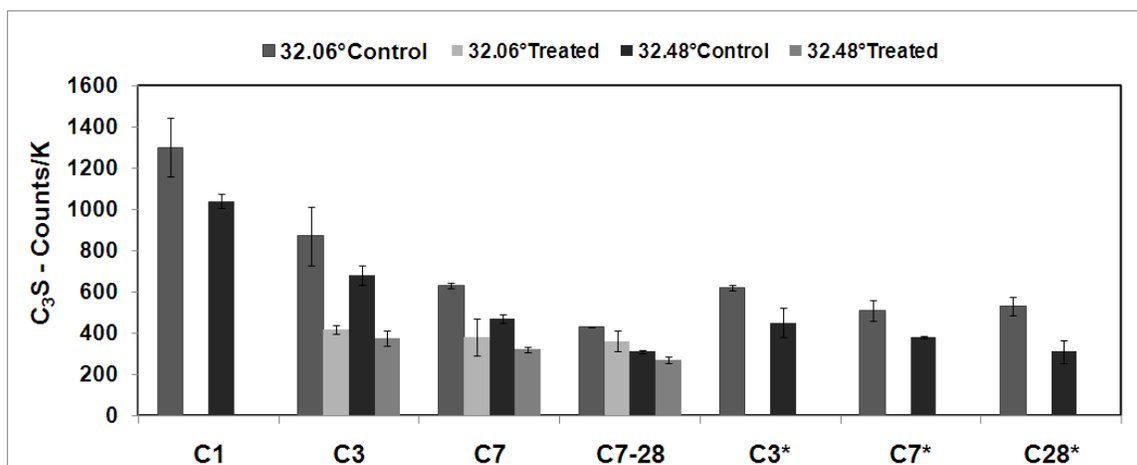


Figure 5

Farahi & Purnell 2012 – Supercritical carbonation of calcareous composite: Influence of curing
Figure 5.

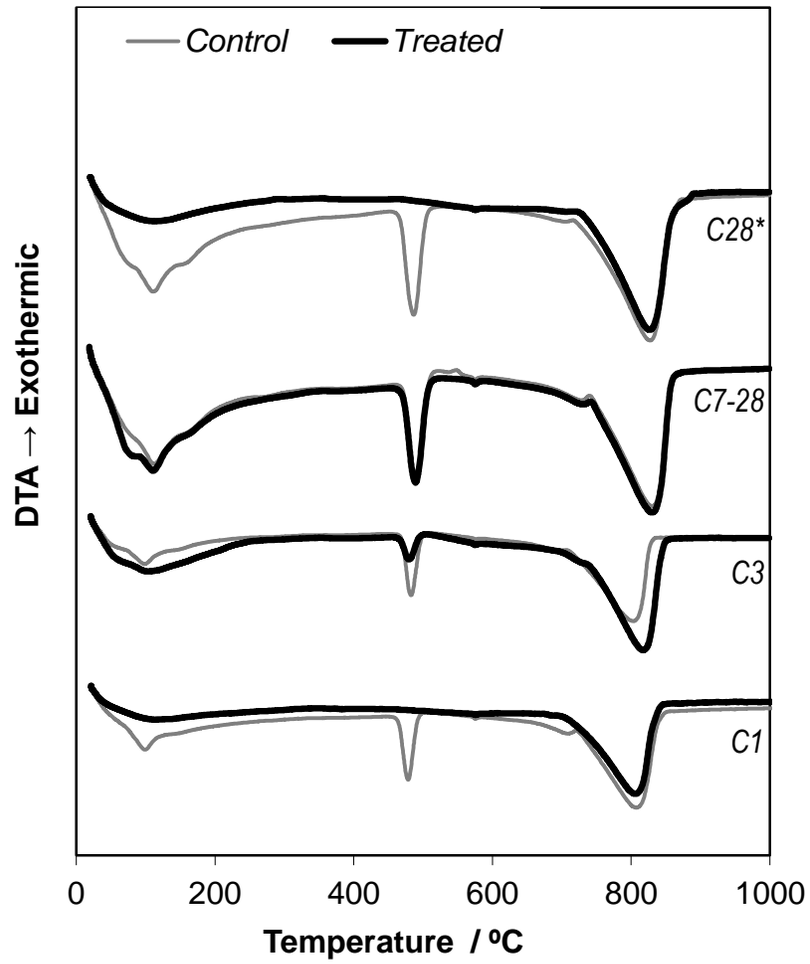


Figure 6

Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing – Figure 6

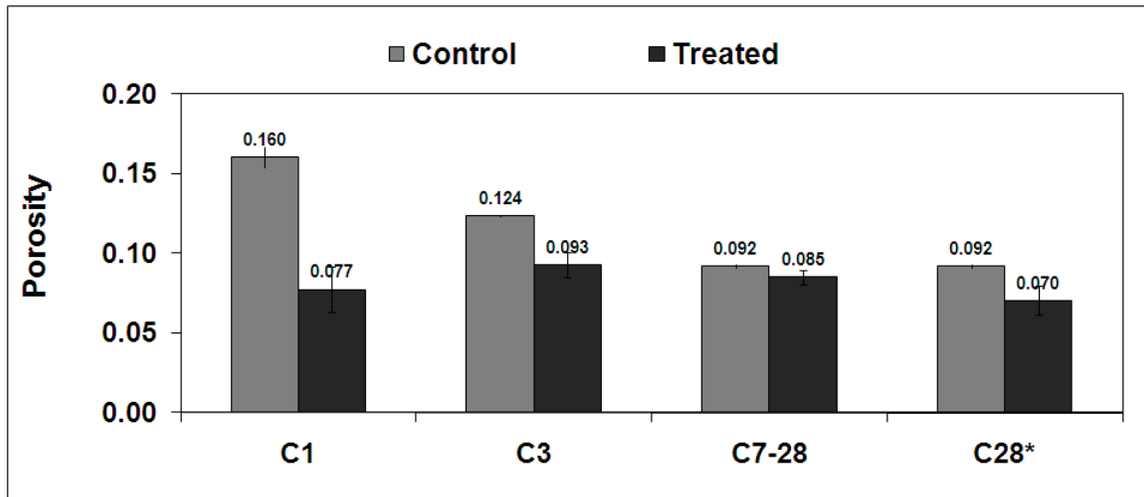


Figure 7

Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing – Figure 7

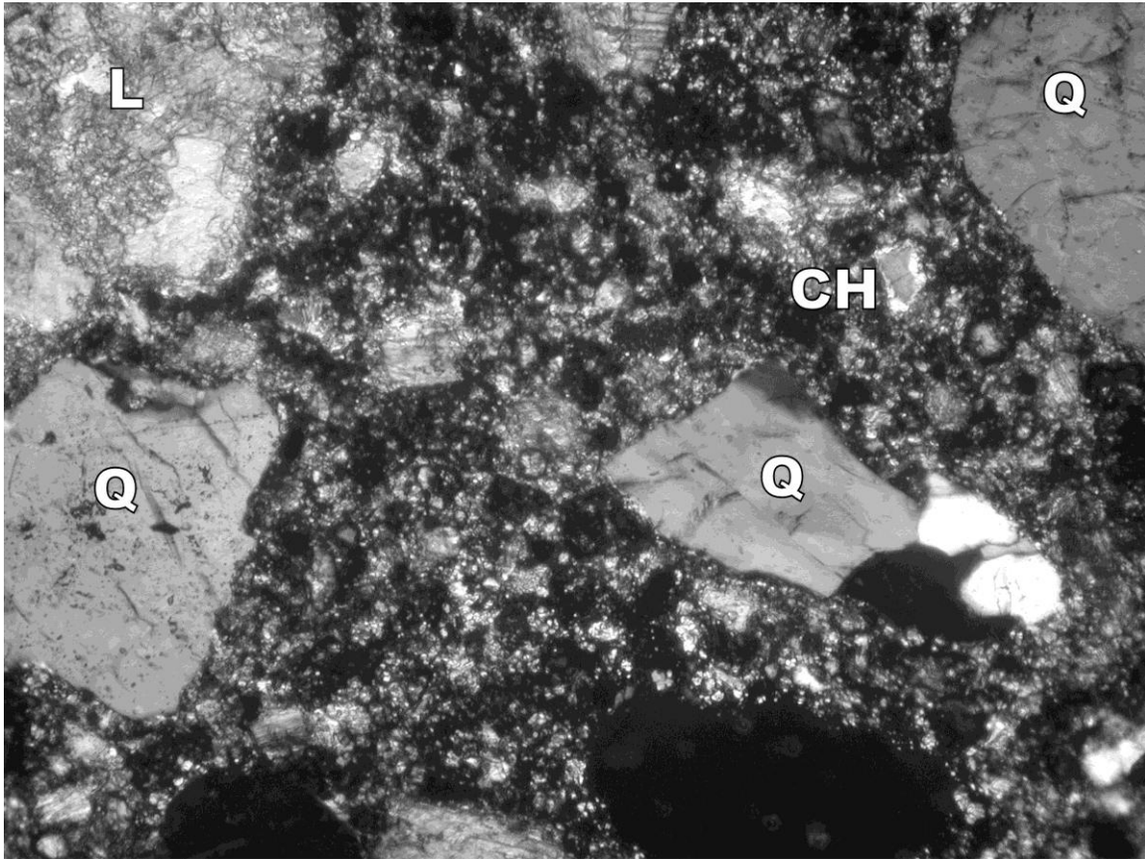


Figure 8

Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing – Figure 8

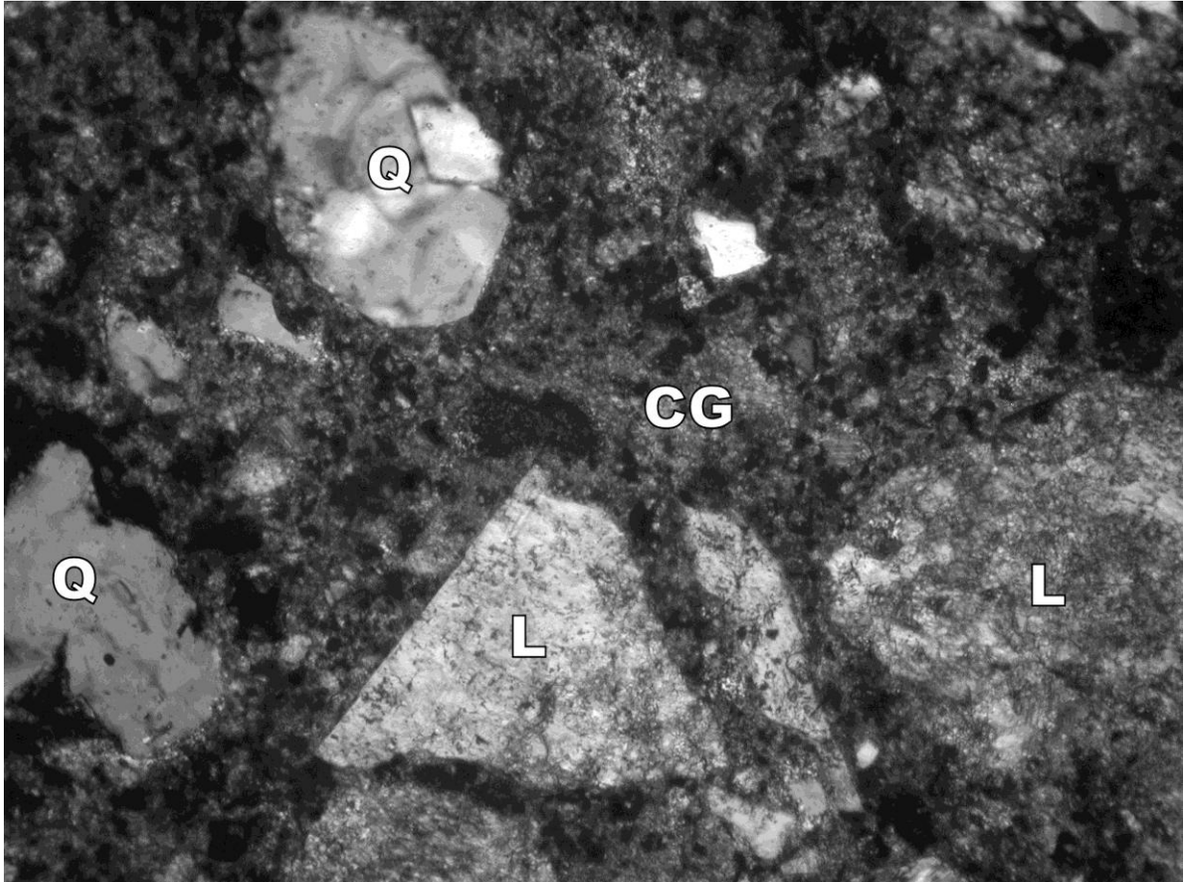
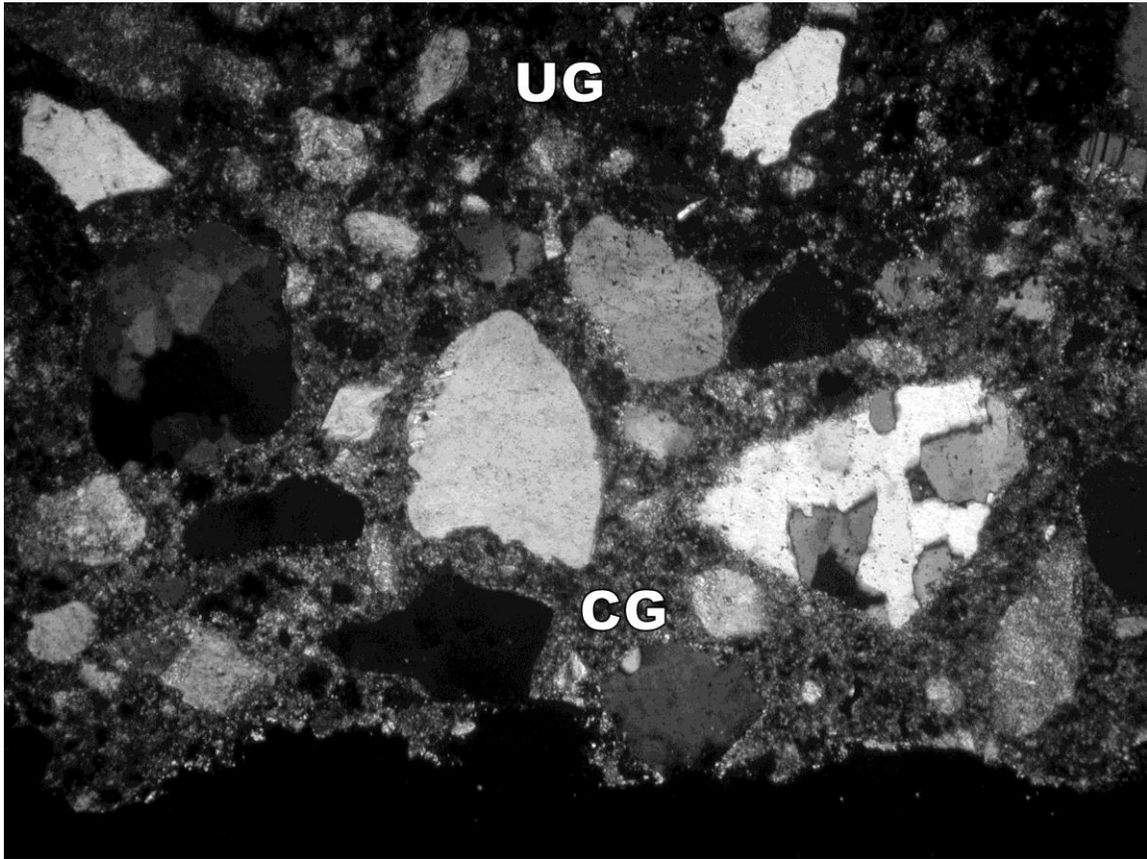


Figure 9

Farahi & Purnell 2012 Supercritical carbonation of calcareous composite: Influence of curing - Figure 9.



Supercritical carbonation of calcareous composite: Influence of curing

Elham Farahi, Phil Purnell, Neil R Short

Figure & table captions.

Figure 1: Layout of the compression moulding tool.

Figure 2: Flexural Strength vs. curing regime. Error bars = \pm st. dev.

Figure 3: XRD analysis of CH depletion vs. curing. Angles refer to 2θ peaks on XRD traces using CuK α radiation wavelength 1.5418 Å. Average of triplicate samples. Error bars = \pm 1 st. dev.

Figure 4: XRD analysis of C3S depletion vs. curing. Angles refer to 2θ peaks on XRD traces using CuK α radiation wavelength 1.5418 Å. Average of triplicate samples. Error bars = \pm 1 st. dev.

Figure 5: DTA thermograms for control and treated samples cured under C1, C3, C7-28, C28* wet curing condition. Black line: Treated samples. Gray Line: Control samples.

Figure 6: Porosity measurements by Helium Pycnometry for mix design 1..5C:0.5L:5A vs. Curing regimes. Treated= 60°C, 100 bar, 24Hr. A =65:56 CL:S. All Proportions w:w. Error bars= \pm st. dev.

Figure 7: Thin section micrograph (TSM) of C1 control sample. Horizontal field of view = 0.7 mm

Figure 8: Thin section micrograph (TSM) of C1 scCO₂-treated sample. Horizontal field of view = 0.7 mm

Figure 9: Thin section micrograph (TSM) of C7-28 scCO₂-treated sample. Horizontal field of view = 1.4 mm

Table 1: Quantitative analysis of TG traces for treated and control samples exposed to following curing regimes. Samples made from; 1.5C:0.5L:5A (A=65:56 CL:S).