Influence of loading and cracks on carbonation of RC elements made of different concrete types

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Abstract

Accurate prediction of concrete carbonation is important for the correct assessment of both durability and environmental impact of reinforced concrete (RC) structures. Loading applied to a RC structure and concrete cracking caused by this loading may significantly affect the concrete carbonation process. However, so far these factors have received little attention of researchers, especially this concerns ‘green’ concretes, i.e. concretes in which Portland cement (PC) is partially replaced by supplementary cementitious materials such as fly ash (FA) and ground granulated blast-furnace slag (GGBS). Thus, the aim of the study presented in the paper was to experimentally investigate the influence of static loading and associated concrete cracking on carbonation of RC elements made of PC concretes and ‘green’ concretes containing significant amounts FA and GGBS. For this purpose, six concrete mixes with two water/binder (w/b) ratios (0.40 and 0.55) and different proportions of PC, FA and GGBS were prepared. The mixes were used to cast twelve RC beams (100×120×900-mm) and a larger number of 100-mm concrete cubes. The beam specimens were loaded in four-point bending to produce flexural cracks of maximum width of either 0.1 mm or 0.3 mm. The loaded beam specimens along with unloaded cube specimens were then placed into a carbonation chamber and subject to accelerated carbonation for 120 days. After that the carbonation depths in the beams and cubes were measured. Results of the tests show a significant effect of load induced stresses (both tensile and compressive) on the carbonation resistance of the concretes, especially of ‘green’ concretes. The influence of cracking on concrete carbonation was also observed and discussed in the paper.

Keywords: accelerated carbonation; Portland cement concrete; fly ash; granulated blast-furnace slag; loading; cracks.

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1. Introduction

Climate change is one of the key challenges facing the human race in the 21st century. This phenomenon is primarily attributed to anthropogenic emissions of greenhouse gases, most notably carbon dioxide (CO₂). One of major global contributors to the anthropogenic CO₂ emissions is the concrete industry, which contributes 5% to those, mainly from a calcination reaction involved in the Portland cement production [1]. CO₂ emissions associated with the concrete industry can be reduced by decreasing the amount of Portland cement used in concrete. This is usually achieved by replacing Portland cement (PC) with supplementary cementitious materials (SCM), mostly industrial by-products such as fly ash (FA), ground granulated blast-furnace slag (GGBS) and silica fume (e.g. [2]) or fillers such as limestone. Concretes containing such materials are often referred to as ‘green’ concretes [3]. A number of studies comparing the environmental impact of traditional and ‘green’ concretes have been published (e.g. [4, 5]).

After a concrete structure has been constructed, the cement paste in the structure becomes exposed to carbon dioxide present in the atmosphere and a carbonation process takes place. During this process, CO₂ diffuses through air-filled pores in the concrete and reacts with the alkaline cement hydration products, in particular with calcium hydroxide, Ca(OH)₂. On one hand, carbonation has a positive environmental effect because CO₂ reabsorbed in this process partially offsets its emission during calcination [6]. On the other hand, the reaction between CO₂ and Ca(OH)₂ reduces the alkalinity of the concrete pore solution and can lead to corrosion of reinforcing steel when the carbonation front reaches the steel (e.g. [7]). This effect of carbonation on the durability of RC structures instigated research of this phenomenon, including experimental studies (e.g. [8-11]), field measurements (e.g. [12-15]) and modelling (e.g. [16-20]). In particular, it has been observed that ‘green’ concretes, in which PC is partially replaced by SCM, can have lower resistance to carbonation than traditional ones (e.g. [10, 21-23]). More recently, the positive effect of carbonation on the CO₂ balance of the concrete industry has also attracted attention of researchers (e.g. [24-26]).

Since natural carbonation (i.e. carbonation under actual atmospheric conditions at which the CO₂ concentration is about 0.04%) is a very slow process, in order to obtain meaningful experimental results
within a reasonable time accelerated carbonation tests with increased CO\(_2\) concentrations are usually performed. The range of the CO\(_2\) concentration used in such tests has varied between 1\% and 100\%, while in most of the tests it has been between 3\% and 10\% [26]. Since an increase in the CO\(_2\) concentration can affect chemical reactions and transport mechanisms involved in carbonation there has been an extensive discussion about the validity of the accelerated tests (e.g. [9, 11, 27, 28]). Although full consensus has not been reached, it has been recommended to keep the partial pressure of CO\(_2\) between 3\%-5\%, in order to obtain results similar to natural conditions; in particular, this has been shown for ‘green’ concretes containing fly ash and blast furnace slag [22, 23]. Among the factors that affect the rate of carbonation, loading and associated cracking have received limited attention and the research has been mainly restricted to PC concretes and effects of tensile stresses caused by static and cyclic loading [29-31]. These studies have demonstrated that tensile stresses/strains have a major effect on carbonation of the elements, i.e. the carbonation rate noticeably increased depending on the stress/strain level [29, 31]. It has also been shown that CO\(_2\) can freely diffuse through cracks with width ≥ 60 μm so that the carbonation depth perpendicular to the crack wall was similar to that at the concrete surface; the diffusion slowed down for smaller crack widths and there was no carbonation penetration perpendicular to the crack wall for crack widths ≤ 9 μm [30]. It is worth to note that the CO\(_2\) concentration used in this study was 50\%.

Thus, the aim of the study described in this paper was to experimentally investigate the influence of static loading and associated cracking on the carbonation performance of RC elements made of PC concretes and ‘green’ concretes containing FA and GGBS. For this purpose, six concrete mixes with two water/binder (w/b) ratios (0.40 and 0.55) and different proportions of PC, FA and GGBS were prepared. The mixes were used to cast twelve RC beams (100×120×900-mm) and 100-mm concrete cubes. The beam specimens were then loaded in four-point bending to produce flexural cracks of different widths. After that both RC beam specimens (loaded) and concrete cubes (unloaded) were transferred to a carbonation chamber. The specimens were subject to accelerated carbonation for 120 days and then the carbonation depths were measured. In the following, the experiments will be described in more detail and
their results presented and discussed. Special attention will be paid to effects on carbonation of both tensile and compressive stresses, concrete cracking and casting position.

2. Experimental Programme

2.1 Materials

Six concrete mixes were used in this experimental study. All mixes included PC (CEM I 52.5 N), in two of them this cement was partially replaced by FA (30%) and in other two by GGBS (50%). The replacement values are typical for ‘green’ concretes. The aggregates in all mixes were Cambusmore sand and 10-mm gravel. Half of the mixes had the w/b ratio of 0.55 and the other half of 0.40; a plasticizer was added to the mixes with the lower w/b ratio. Table 1 shows the mix constituent proportions per cubic metre of concrete and was provided by the precaster, who produced the test specimens prior to transferring them to the laboratory. 8 mm and 10 mm reinforcing bars used in the specimens had yield strength of 500 MPa.

2.2 Test specimens

Twelve RC beams with a rectangular cross section of 100×120 mm and length 900 mm were cast. The beams were reinforced in the longitudinal direction with two 10-mm diameter deformed bars at the bottom (tensile reinforcement) and two 8-mm diameter bars at the top (compression reinforcement), and in the transverse direction with stirrups made of 8-mm diameter bars. The concrete cover of the longitudinal tensile bars was to be 25 mm. The spacing between the stirrups was 46 mm near the beam ends and 105 mm closer to the midspan, while the central 200-mm long part of the beams was without shear reinforcement. Two vertical (i.e. along the beam depth) 18-mm diameter holes at 100 mm from the beam ends (so that the distance between the holes was 700 mm) were made in each beam in order to create later a four-point bending system. Figure 1 shows the beam dimensions and reinforcement details. The beams were cast horizontally in wood moulds. In addition to the beams, seven 100-mm cubes were cast from each mix to determine the compressive strength and porosity of concretes made of the mixes and also the carbonation rate of unloaded specimens. All specimens were standard cured for 28 days.

As noted previously, the specimens were fabricated by a precaster. Thus, compared to the fabrication
of specimens in a laboratory, the composition and quality of the specimens were not fully controlled by the researchers. However, the tests were probably more realistic since the specimens were more representative of typical RC elements produced by the industry.

2.3 Preconditioning of the specimens

All specimens (i.e. both beams and cubes) intended for accelerated carbonation were kept for 3 months under ambient conditions (i.e. temperature and RH) similar to those in the carbonation chamber. The beam specimens were then quasi-statically loaded in four-point bending using a universal testing machine, as shown in Figure 2. In accordance to the beam dimensions (Figure 1), the distance between two supporting pins was 700 mm (i.e. they were placed at the location of the holes) and the distance between two loading pins 200 mm. The level of the applied load was controlled to ensure that the maximum width of cracks in the tensile zone of the beams due to bending was either 0.1 mm or 0.3 mm. The load required to induce 0.1-mm wide cracks was about two-thirds of that used to induce 0.3-mm wide cracks. At the same time, three concrete cubes from each mix were tested to determine the compressive strength of the concretes.

After the beams had been loaded and cracked, they were removed from the testing machine and placed in pairs, the top surface of one beam to the top surface of the other, to create four-point bending like in the initial loading state. This time two pins of 25-mm in diameter were placed at a distance of 200 mm from each other between the beams in each pair (at the previous locations of the loading pins), while two 16-mm diameter steel bars were put through the holes in the beams and tightened until the cracks in the beams opened to approximately the same widths as previously in the testing machine. The pre-tensioned bars were then locked in place (Figure 3) and the beam pairs along with the cubes were placed into the carbonation chamber.

The actual dimensions of the beams and cracks in each beam had been measured and recorded before the beams were put into the carbonation chamber (see Table 2). The beam designation identifies the concrete mix followed by two digits that define the maximum width of the initial flexural cracks (e.g. ‘040PC01’ is a PC concrete beam with a w/b ratio of 0.40 and the maximum initial crack width of 0.1 mm). It should be noted that the beams were placed in pairs based on the concrete properties and not on the initial
crack widths, i.e. all pairs except of one consisted of beams made from the same concrete type, e.g. 040PC01 and 040PC03, 055PC01 and 055PC03, etc. The only exception was a pair consisting of 040FA03(2) and 040GGBS03 (the second beam made from the 040GGBS concrete was not properly fabricated and had to be replaced by a spare beam from 040FA). This led to noticeable differences between the maximum crack widths measured after the beams were coupled in pairs (Table 2) and the maximum initial crack widths of 0.1 mm and 0.3 mm obtained in the testing machine.

2.4 Accelerated carbonation test and carbonation depth measurements

The beam specimens and the concrete cubes were kept in the carbonation chamber for 120 days. The chamber conditions were as follows: T=20±5°C, RH=60±5% and CO₂ content of 4±0.50%. The specimens were not removed from the chamber for the whole duration of the test so that the tensile forces in the pre-tensioned bars in the beam pairs could be partially lost over the time due to creep. The duration of the test was selected to ensure sufficiently large carbonation depths in most of the test specimens. This should reduce uncertainties associated with the carbonation depth measurements, which can be significant when the carbonation depth is small [32].

After the specimens had been removed from the carbonation chamber they needed to be split in halves to measure the carbonation depth. To do that the central parts of the beams (about 500 mm long), which had been under load during the accelerated carbonation test, were cut along their length into five segments. The segments were then split or diamond sawn (if they contained stirrups) along their lengths. The split/sawn surfaces were sprayed with a phenolphthalein solution (see Figure 4) and the carbonation depths measured at 10 mm intervals. Additional measurements were taken at the locations where changes of the carbonation depth were observed (e.g. around cracks). The concrete cubes were tested in accordance to BS EN 12390-7:2009 [33] to determine the absorption and void content of the concretes as shown in Table 3.

3. Test Results

3.1 Concrete properties

As these mixes used direct replacement of PC with SCM, the 040PC concrete had the highest
compressive strength and the 055FA – the lowest one (Table 2). However, the GGBS concretes had compressive strengths close to those of the PC concretes and for the w/b ratio of 0.55 the compressive strength of the 055GGBS was even higher than that of the 055PC. The latest result is in line with the values of the void content shown in Table 3. As has been shown by previous research, lower porosity typically corresponds to higher compressive strength [34, 35]. In fact, this trend can be observed for practically all values of the compressive strength and void content presented in Tables 2 and 3. The only exception is the 040GGBS concrete, which had slightly lower porosity than the 040PC concrete but also lower compressive strength. This seemed unexpected that the GGBS concretes had lower porosity than the PC ones even though the reactions associated with GGBS had a slower rate compared to the hydration of PC. However, it is known that adding GGBS to PC concretes eventually leads to reduced porosity [36]. Since the void content in the tests was measured in about 8 months after casting the specimens there was enough time to achieve this effect. It should also be noted that values of the compressive strength given in Table 2 were obtained after four months and not on 28 days. The older age of the concretes at the time of the compressive tests probably explains the small differences between the measured compressive strengths of the PC and GGBS concretes. It is worth to note that the main characteristic of concretes used in this study is the w/b ratio. This is in line with the fib Model Code 2010 [37], in which the carbonation resistance of concretes is also primarily related to their w/b ratio.

3.2 Cracks in RC beam specimens

Cracks in the beam specimens were measured and recorded right before placing the specimens into the carbonation chamber. Three types of cracks were observed: (1) flexural cracks in the tensile zone at the bottom of the specimens; (2) shear cracks in the zones with shear stresses (i.e. zones within the loaded part of the beams outside the 200-mm central part); and (3) drying shrinkage cracks at the top of the beams (i.e. near the casting surface). The recorded crack patterns on the lateral sides of the beam specimens are shown in Figures 6 – 8. For each beam specimen two crack patterns (i.e. on both lateral sides of the beam) were recorded and the records were distinguished as “L – R” and “R – L”, i.e. from the left end of the beam to its right end and vice versa for the opposite side. Since most cracks extended through the whole beam width
the patterns on the two lateral sides were usually quite similar so that only one of them is shown in the figures. The crack widths given in the figures were recorded at the bottom of the beams; their maximum values for each beam are also given in Table 2.

The specimens with the initial crack widths of 0.3 mm usually had wider cracks when loaded in the pairs than the ones with the initial 0.1-mm wide cracks. Moreover, the specimens with the initial 0.3-mm wide cracks also had noticeable shear cracks (e.g. 055PC03, 040FA03, 055FA03, etc.), while the ones with the initial 0.1-mm cracks either had none (e.g. 055PC01) or very few (e.g. 040FA01, 055FA01, etc.), see Figures 5 – 7. This can be explained by higher loads that had to be applied to the specimens in the universal testing machine to obtain wider flexural cracks. All specimens had hairline drying shrinkage cracks near the top (casting) surface. However, in the specimens made from the PC concretes these cracks were shallow, while in the specimens made from the FA and GGBS concretes the drying shrinkage cracks were deeper and nearly connected with flexural cracks, so that the cracks practically extended through the whole beam depth (e.g. 040FA03, 055FA01, 040SG03, 055SG01).

As noted previously, the specimens were not removed from the carbonation chamber for the whole duration of the carbonation test (120 days) and loads in the pairs were not checked over that time. Thus, the loads could decrease due to creep and further shrinkage strains and the cracks partially close (compared to their dimensions shown in Figures 5 – 7) while the specimens were in the carbonation chamber.

3.3 Carbonation depths

As explained above, two types of specimens were used in the test: unloaded 100-mm concrete cubes and loaded RC beams. To measure carbonation depths in the cubes the latter were split in the middle into halves and the depths were measured for each half. The average carbonation depths at the top (cast surface) and bottom of the cubes are shown in Table 4. The influence of two-dimensional CO₂ penetration on the carbonation depth was clearly visible near the corners of the specimens. Thus, the carbonation penetrations near the corners were not taken into account in the calculations of the average carbonation depths. For all cubes the average carbonation depth at the top (i.e. near the cast surface) was larger than that at the bottom, which is typical and due to free water migration towards the as-cast surface.
The measured carbonation profiles for the beams are shown in Figures 8 – 10. They are shown for the same halves of the beams as the crack patterns in Figures 5 – 7. However, the crack patterns were recorded on the external lateral surfaces of the beams, while the carbonation depths were measured on the internal surfaces of the halves. Thus, an “R – L” carbonation profile corresponds to the “L – R” crack pattern with the same name and vice versa. In these figures, the measured carbonation profiles are shown by black lines, vertical carbonation depths at crack locations (i.e. flexural cracks at the bottom and drying shrinkage cracks at the top) are indicated by red numbers and red straight lines with arrows at both ends, and horizontal carbonation penetration through the crack walls are indicated by blue numbers and blue straight lines with arrows at both ends. Some visible cracks are also shown by red lines.

Results of the carbonation depth measurements in the beam specimens are summarised in Table 4. The average carbonation depths were calculated based only on the measurements within the central 200-mm long parts of the beams, within which the bending stresses (i.e. tensile at the bottom and compressive at the top) were constant. This was done in order to exclude the influence of the stress magnitude on the carbonation depth for the same specimen, as noted in previous research [29]. Moreover, the influence of visible flexural cracks has also been excluded, i.e. the carbonation depths measured within the vicinity of visible flexural cracks have not been taken into account in the calculations of the average carbonation depths. The maximum carbonation depths given in Table 4 are based on all measurements for a particular specimen and usually coincide with the location of major flexural cracks. The only exception was the 055FA01 specimen. As can be seen in Figure 9(c), the maximum carbonation depth (75.9 mm) was measured at the end of the central segment cut out from this specimen and there is a clear discontinuity in the carbonation depth between this segment and the adjacent one. This occurred due to the central segment being split unevenly along its length and the width of one of the ‘halves’ near its end being smaller than the half-width of the beam. Thus, the measured carbonation depth of 75.9 mm does not represent the carbonation penetration through the bottom surface of the beam (which is of interest) but is rather due to the carbonation penetration through the nearest lateral beam surface. Hence, the carbonation measurements near this end of the central segment have been excluded from the analysis of the test results.
Carbonation penetration for the 040PC03 specimen was observed only in a couple of isolated locations and thus was not included in the analysis.

4. Discussion

4.1 Influence of concrete type and water-binder ratio on carbonation resistance

The resistance of concretes to carbonation in the context of their properties depends mainly on their CO$_2$ binding capacity and degree of interconnected porosity (e.g. [38]). An increase in the w/b ratio increases the concrete porosity and, subsequently, decreases the concrete resistance to carbonation. Results of the tests for the same concrete type (i.e. PC, FA or GGBS) follow this trend. As can be seen from Table 3, the porosity of concretes of the same type with the w/b ratio of 0.55 is larger than those with w/b=0.40 and the average carbonation depth in the cubes is correspondingly larger (see Table 4). The situation becomes more complicated when PC concretes are compared with concretes with SMC and concrete elements are subject to loading. The latter will be discussed in the next section.

Replacement of PC in concrete with SCM reduces the amount of carbonatable constituents and, subsequently, the CO$_2$ binding capacity of the concrete that should lead to an increase in the carbonation rate. Direct replacement mix designs usually result in higher porosity of the concrete when the w/b ratio is kept constant, although, as noted previously, concretes containing GGBS may eventually have lower porosity than PC ones with the same w/b ratio. In these tests, the 040PC concrete demonstrated the highest carbonation resistance although its porosity was slightly higher than that of the 040GGBS concrete. However, this was different for the PC and GGBS concretes with the w/b ratio of 0.55 – where the PC concrete had higher porosity than the 055GGBS (see Table 3), while its carbonation resistance was lower (see the corresponding average carbonation depths for the cubes in Table 4). This data shows that the lower porosity had a more significant effect on the carbonation resistance of the 055GGBS concrete than its reduced CO$_2$ binding capacity compared to the 055PC.

The 040FA concrete also had lower porosity than the 055PC and 055GGBS concretes (see Table 3) but its carbonation resistance was also lower. The 055FA concrete had the highest porosity and the lowest carbonation resistance in unloaded state (see Table 4).
Based on the average carbonation depths for the cubes (Table 4), the concretes considered in this study can be ranked in the following order in terms of their carbonation resistance in unloaded state (from lowest to highest): 055FA, 040FA, 055PC, 055GGBS, 040GGBS and 040PC. It is important to highlight that these mixes did not have equal strength and that the rank order is likely to be different when based on comparative strength rather than w/b.

The results for the FA concretes showing their lower carbonation resistance compared to that of PC and GGBS concretes are in line with most of the previous studies [22, 23]. There is less consensus regarding the relative carbonation resistance of GGBS and PC concretes. It is often suggested that the carbonation resistance of GGBS concretes is lower than that of PC concretes with a similar w/b ratio, especially when the GGBS content is greater than 20% [23]. However, according to several studies, concretes with 50% GGBS content have more or less the same carbonation resistance as PC concretes with a similar w/b ratio [39, 40] or similar strength grade [41, 42]. The results of this study for GGBS and PC concretes (with w/b=0.55) in unloaded state support the latter observations.

4.2 Influence of loading on carbonation resistance

Since in this study both unloaded and loaded (in four-point bending) specimens were subject to accelerated carbonation testing this allows to assess the influence of both compressive and tensile stresses in concrete on its carbonation. First of all, it is worth to note that while in each tested concrete cube the average carbonation depth at the top was larger than that at the bottom (due to casting direction) the opposite was observed in the beam specimens although the casting position was the same, i.e. the top was the beams’ casting surface (see Table 4). This demonstrates that compressive and tensile stresses had more significant influence on the carbonation resistance than the casting direction.

The results in Table 4 also show that tensile stresses led to a decrease in the carbonation resistance, since the average carbonation depth at the bottom of each beam specimen was larger than that of the corresponding concrete cube. This is in agreement with previously reported experimental results [29]. As can be seen, the largest increase in the average carbonation depth due to tensile stresses was observed in the beams made of the concretes with SCM and the w/b ratio of 0.55, i.e. 055FA (48.1 mm in 055FA01 and 38.8
mm in 055FA03 versus 18.5 mm in the corresponding cube) and 055GGBS (24.4 mm and 29.8 mm in the beams versus 12.0 mm in the cube). Regarding the beams made from 055FA, it is necessary to note that the results could be partially influenced by flexural cracks; in particular, for the 055FA01, in which the flexural cracks were spaced much closer (average spacing 60 mm – see Figure 6(c)) than in the other specimens (average spacing 90-95 mm – see Figures 5-7). Thus, it was difficult to distinguish between the vertical penetration of CO₂ through the bottom surface of the 055FA01 beam and the horizontal penetration through the crack walls. In the unloaded state, the 055GGBS concrete had a higher carbonation resistance than the 055PC and 040FA mixes due to its lower porosity than that of 055PC and higher CO₂ binding capacity than that of 040FA. Damage to the concrete (i.e. microcracking) caused by tensile stresses negated these advantages of the 055GGBS in the context of carbonation resistance - as can be seen from Table 4 the average carbonation depth at the bottom of the 055GGBS beams is larger than that at the bottom of the 055PC and 040FA beams. As in the unloaded state, under tensile stresses the 055FA concrete had the lowest carbonation resistance among all considered concretes. According to the average carbonation depths at the bottom of the beam specimens shown in Table 4, the concretes can be ranked in the following order in terms of their carbonation resistance under tensile stresses (from lowest to highest): 055FA, 055GGBS, 040FA, 055PC, 040GGBS and 040PC. Again, it is important to note the caveat that this is not based on equal strength.

While in the past the main attention has been on the influence of tensile stresses on the carbonation resistance of concrete (e.g. [29, 31]) in this research the influence of compressive stresses has also been studied. Based on the average carbonation depths measured at the top (i.e. in compression zone) of the beam specimens (Table 4), compressive stresses have a positive effect on the carbonation resistance of concrete. One possibility to explain this is that compressive stress closes microcracking and hence reduces the degree of interconnected voidage but since this could only be measured on unloaded specimens it was not possible to confirm this hypothesis. Only in three beam specimens: 055FA01, 055GGBS01 and 055GGBS03, the average carbonation depth at the top was larger than that at the top of the corresponding concrete cubes. However, the presence of comparatively large drying shrinkage cracks in these three beams
(see Figures 6(c), 7(b) and (c)) potentially negated the effect of compressive stresses on microcracks. Based on the average carbonation depths for the beams made from the same mix type, the concretes can be ranked in the following order in terms of their carbonation resistance under compressive stresses (from lowest to highest): 055FA, 055GGBS, 040FA, 055PC, 040GGBS and 040PC, which is the same that noted for the tensile stressed concrete.

Thus, comparing the results for the loaded and unloaded states it can be concluded that the 055GGBS concrete has been more negatively affected by loading, in terms of the carbonation resistance, than the 055PC and 040FA concretes. The positive effect of GGBS on the concrete porosity (and, subsequently, on the carbonation resistance) was negated by microcracks due to tensile stresses in the tension zone and drying shrinkage cracks in the compression zone. The other concrete, where carbonation resistance was significantly affected by loading, was the 055FA (see Table 4). However, this mix demonstrated the lowest carbonation resistance in both loaded and unloaded states.

4.3 Influence of cracking and other factors on carbonation resistance

As noted previously, three types of cracks: flexural, shear and drying shrinkage, were observed in the beam specimens. The influence of flexural cracks on the vertical carbonation penetration (i.e. through the bottom surface of the beams) can be observed in all beam specimens – the maximum carbonation depths at the bottom of the beams, which are summarised in Table 4, coincide with the location of the flexural cracks. The only exception is the 055FA01 beam, in which the maximum carbonation depth (75.9 mm) according to the carbonation profile shown in Figure 9(c) is on the border between two segments. However, as explained previously in Section 3.3, this result does not represent the CO₂ penetration through the bottom surface of the beam and should be excluded from consideration. The next largest value (64.1 mm), which is given in Table 4, is at the location of a flexural crack. There is also carbonation penetration perpendicular to the crack walls, which according to previous research depends on the crack width. In particular, it was observed that in cracks of 0.06-mm wide or wider the depth of this penetration was similar to that at the concrete surface [30]. The CO₂ penetration perpendicular the walls of the flexural cracks can be clearly seen in a number of the carbonation profiles measured in the present study,
e.g. 055PC01 (Figure 8(a)), 040FA01 (Figure 9(a)), 040FA03(2) (Figure 9(b)), 040GGBS03 (Figure 10(a)) and 055GGBS03 (Figure 10(c)). At the same time, it can also be seen that no (or almost no) CO2 penetration perpendicular to the crack wall was observed for many flexural cracks appearing in the crack patterns (Figures 5-7). As noted previously, the crack patterns had been recorded before the beams were placed into the carbonation chamber. Since loads applied to the beams were not controlled while the beams were in the carbonation chamber the loads could decrease due to concrete creep and the cracks could partially (or fully) close. Alahmad et al. [30] noted that no CO2 penetration perpendicular to the crack wall was observed for cracks narrower than 0.01 mm; however, the CO2 concentration used in their tests was 50%. Thus, it can be expected that in the present tests with the CO2 concentration of 4% no noticeable CO2 penetration perpendicular to the crack wall was observed even for slightly wider cracks.

The 055FA01 beam is a special case – as the spacing between flexural cracks in this beam was smaller than in the other beams (on average 60 mm versus 90-95 mm in the others) so that areas between the flexural cracks could be fully carbonated due to CO2 penetration through the crack walls that created an impression of more or less uniform vertical propagation (i.e. from the bottom of the beam upwards) of the carbonation front (see Figure 10(c)).

Shear cracks were observed in the beam specimens that were initially loaded to create 0.3 mm flexural cracks. Among these specimens, the CO2 penetration perpendicular to the shear crack walls can be seen in 055PC03, 055FA03 and 055GGBS03 (Figures 8(b), 9(d) and 10(c), respectively). Larger opening of the shear cracks in these specimens that caused this penetration can be explained by lower strength of the specimens’ concretes compared to the corresponding 0.40 w/b specimens.

Drying shrinkage cracks had a smaller effect on the carbonation propagation compared to the flexural cracks, e.g. the maximum carbonation depths at the top (see Table 4) rarely coincide with the location of drying shrinkage cracks (see Figures 8-10). This is probably due to compressive stresses reducing the crack widths. In fact, in most specimens the average carbonation depth at the top (that was calculated taking into account all carbonation measurements there) is less than that in the corresponding unloaded cubes (see Table 4), which did not have drying shrinkage cracks. The exception are the beams
made from the 055GGBS concrete – in these two beams the average carbonation depths at the top are larger than in the corresponding cube that can be attributed to influence of the drying shrinkage cracks.

Another factor that influenced the carbonation of the beam specimens is the use of stirrups. It has been shown previously that carbonation occurs at the steel-concrete interface, where it intercepts a crack, irrespective of the crack width [30, 43]. The same phenomenon was observed in these tests, e.g. larger carbonation depths under stirrups at the interception of drying shrinkage cracks (i.e. at the top of the beam) are clearly noticeable in, e.g., 050PC01 (Figure 8(a)), 050PC03 (Figure 8(b)), 055FA03 (Figure 9(d)) and 040GGBS03 (Figure 10(a)).

5. Conclusions

Accelerated carbonation tests of PC and ‘green’ (with FA and GGBS) concretes in unloaded and loaded states have been described. The emphasis of this research has been on studying the influence of loading, i.e. imposed stresses (both tensile and compressive), cracking and casting position on the carbonation of the concretes.

Based on the data generated the following conclusions can be made:

1) The concrete composition, including the w/b ratio, had a major influence on the carbonation resistance of the concretes. As expected, concretes of the same type with the larger w/b ratio of 0.55 had lower carbonation resistance than those with the w/b ratio of 0.40. Generally, replacing PC with FA and GGBS led to a reduction of the carbonation resistance. The only exception is the 055GGBS concrete that had higher carbonation resistance in the unloaded state than the 055PC concrete. The concretes with FA had lower carbonation resistance than those with PC and GGBS with the same w/b ratio. Based on the w/b ratio the concretes can be ranked in the following order in terms of their carbonation resistance in the unloaded state (from lowest to highest): 055FA, 040FA, 055PC, 055GGBS, 040GGBS and 040PC.

2) The casting position had a noticeable influence on the carbonation resistance of the concretes in the unloaded state. In all unloaded specimens the carbonation depth at the top (i.e. casting surface) was larger than that at the bottom. This can be attributed to better concrete compaction at the bottom.

3) Loading had a major influence on the carbonation resistance of the concretes. Generally, compressive
stresses led to an increase in the carbonation resistance of the concretes, while tensile stresses resulted in its decrease. The 055GGBS and 055FA concretes were mostly affected by loading in terms of their carbonation resistance. The concretes can be ranked in the following order in terms of their carbonation resistance under loading (from lowest to highest): 055FA, 055GGBS, 040FA, 055PC, 040GGBS and 040PC.

4) Flexural cracks clearly affected the upward propagation of the carbonation front. In all beam specimens the maximum carbonation depths at the bottom coincide with the location of the flexural cracks. The penetration of CO₂ perpendicular to the walls of flexural and shear cracks was also observed. In a number of the beam specimens the depths of this penetration were similar to those at the beam surface. Drying shrinkage cracks did not have such a noticeable effect on the carbonation propagation as flexural cracks. It can be attributed to the location of the drying shrinkage cracks in the compression zone of the beams so that compressive stresses caused the cracks’ closure. The only exception is the beams made from the 055GGBS concrete, in which the average carbonation depth at the top was larger than that in the corresponding unloaded specimen. Thus, multiple drying shrinkage cracks in these beams negated the positive effect of compressive stresses so that the carbonation resistance of the concrete at the top decreased compared to the unloaded state.

Acknowledgments

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References


[28] J.H.M. Visser, Influence of the carbon dioxide concentration on the resistance to carbonation of...


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Figure 3. Beam specimens in pairs – setup for four-point bending in carbonation chamber.

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Figure 5. Crack patterns on lateral surfaces of the beam specimens made from 055PC concrete: (a) 055PC01; (b) 055PC03.

Figure 6. Crack patterns on lateral surfaces of the beam specimens made from FA concretes: (a) 040FA01; (b) 040FA03(2); (c) 055FA01; (d) 055FA03.

Figure 7. Crack patterns on lateral surfaces of the beam specimens made from GGBS concretes: (a) 040GGBS03; (b) 055GGBS01; (d) 055GGBS03.

Figure 8. Carbonation profiles at the middle width of the beam specimens made from 055PC concrete: (a) 055PC01; (b) 055PC03.

Figure 9. Carbonation profiles at the middle width of the beam specimens made from FA concretes: (a) 040FA01; (b) 040FA03(2); (c) 055FA01; (d) 055FA03.

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(a)

Drying shrinkage cracks

(b)
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Drying shrinkage cracks

Average carbonation depth within the zone = 10.6 mm

Uncarbonated zone

Uncarbonated zone

Uncarbonated zone

Uncarbonated zone

Uncarbonated zone

Pure bending moment zone

Average carbonation depth within the zone without influence of visible flexural cracks = 1.5 mm

0.12 mm crack 0.08 mm crack 0.34 mm crack 0.4 mm crack 0.42 mm crack 0.24 mm crack

Drying shrinkage cracks

Average carbonation depth within the zone = 21.6 mm

Uncarbonated zone

Uncarbonated zone

Uncarbonated zone

Uncarbonated zone

Uncarbonated zone

Pure bending moment zone

Average carbonation depth within the zone without influence of visible flexural cracks = 24.4 mm

0.2 mm crack 0.22 mm crack 0.22 mm crack 0.3 mm crack 0.1 mm crack 0.06 mm crack
Figure 10. Carbonation profiles at the middle width of the beam specimens made from GGBS concretes: (a) 040GGBS03; (b) 055GGBS01; (c) 055GGBS03.
### Table 1 Mix constituent proportions of concretes (kg/m³)

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<th>Sand</th>
<th>10-mm gravel</th>
<th>Plasticizer</th>
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### Table 2 Details of RC beam specimens

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<th>Test specimens</th>
<th>Cross section (mm)</th>
<th>Average clear concrete cover of 10-mm bars in tensile zone (mm)</th>
<th>Average clear concrete cover of stirrups</th>
<th>Maximum width of flexural cracks (mm)</th>
<th>Concrete compressive strength (MPa)</th>
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Table 3 Absorption and void content of concretes

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*The mass includes the mass of the collar (66.6 g) that supported the immersed concrete cube in water.
Table 4 Carbonation depths in specimens

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*Over the 200-mm long central segment excluding the influence of visible flexural cracks*