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Removal of Hexavalent Chromium in Portland Cement Using Ground Granulated Blast-Furnace Slag Powder

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Abstract: Using ground granulated blast-furnace slag (GGBS) under different alkaline conditions, we studied the mechanisms and extents of Cr(VI) reduction and sorption and compared them to reactions with Portland cement (PC). We also investigated the effects of mixing PC/GGBS ratios on Cr(VI) dissolution after carbonating the substrates. We observed a complete sorption and reduction of Cr(VI) to Cr(III) in a GGBS-in-Ca(OH)2 solution (pH > ~12.5) after 10 h, whereas in distilled water (pH ~11.5) GGBS exhibited only marginal sorption and reduction (20%). Cr reactions with dissolved ions in supernatants derived from GGBS indicated that the anions dissolved from GGBS act as a reducing agent for Cr(VI) in a Ca(OH)2 solution. Soft X-ray absorption microscopy identified a partial reduction of Cr(VI) to Cr(III) on the GGBS surface. The carbonation of pure PC paste substantially increased the amount of dissolved Cr(VI) in a solution phase whereas a 5 wt % replacement of PC with GGBS significantly reduced the amount of dissolved Cr(VI). We concluded that in the mixed paste during the early curing stage GGBS reduced a significant fraction of Cr(VI) to Cr(III) and that the Cr(III) adsorbed in the GGBS-PC mixture’s hydration products does not readily dissolve, even under carbonation conditions.

Keywords: hexavalent chromium; reduction; immobilization; slag; Portland cement

1. Introduction

Hexavalent chromium [Cr(VI)] is found in concrete as chromate, which results from the oxidation process of trivalent chromium [Cr(III)] during Portland cement (PC) clinker production [1,2]. Chromium (Cr) can exist in several different oxidative states and its physical and chemical properties, such as solubility, toxicity, and sorption affinity, vary with environmental conditions. For example, Cr(III) is likely to precipitate in alkaline solutions, whereas Cr(VI) is amphoteric with a concentration-dependent solubility [3]. Cr(VI) is highly mobile but very toxic under aerobic conditions [1,4,5]. Cr(III) has a strong affinity to soils and other substrates; while it is less soluble and toxic than Cr(VI), its potential for oxidation to Cr(VI) should be carefully examined in order to fully understand Cr transport and distribution in the environment. Therefore, examining Cr(VI) dissolution,
sorption, and reduction in PC clinker is of the utmost important for environmental concerns and for the long-term sustainability of PC materials in construction materials [3,6].

In the past, stabilization and solidification (s/s) have been practiced on immobilized PC-associated Cr(VI) [7–9]. Stabilization involves a reductive transformation of Cr(VI) to Cr(III) and a subsequent chemical transformation into a highly insoluble compound, such as calcium chromate [10,11]. Solidification involves the co-precipitation of Cr(VI) with dissolved PC hydration products during hydration reactions [10]. Calcium silicate hydrate (C-S-H), which is the primary hydration and binding product in PC-based materials, contains most of the microporosity, resulting in a large specific surface area that provides physical sorption and co-precipitation sites for heavy metal ions like Cr(VI). It should also be noted that Zamorani et al. [12] observed co-precipitated phase of CaCrO$_4$ when Cr(VI) reacted with C-S-H, suggesting that molecular-level surface reactions are important in determining the reaction rates and pathways when hydrated products of PC-based materials react with Cr(VI) [3]. Ettringite (3CaO·Al$_2$O$_3$·3CaSO$_4$·32H$_2$O) has also been identified as one of PC’s hydration products and, with its highly reactive and needle-like column structure, has been proposed as a surface site for Cr(VI) immobilization via anion substitution, where CrO$_4^{2−}$ is exchanged for SO$_4^{2−}$ [13,14]. Another study identified that reductive reactions occur during the curing phases and that the resulting Cr(III) also replaces Al(III) in the ettringite crystalline structure [15]. Ionic substitutions in ettringite were also observed to be Cr(VI)–ettringite and Cr(III)–ettringite phases that further limit the mobility of Cr [8,16]. Cr(VI) can be substituted for SO$_4^{2−}$ in the interlayer of calcium aluminate monosulfate (hereafter referred to as monosulfate, 3CaO·Al$_2$O$_3$·CaSO$_4$·12H$_2$O) [16]. However, the pH stability range of reactions for Cr(VI)–ettringite and Cr(VI)-hydration products of PC is very narrow and varies significantly [16,17].

The alkalinity of the PC-based matrix is important for decreasing the dissolution of Cr ions [3,11]. However, the carbonation process neutralizes the alkaline conditions and increases the Cr(VI) solubility for PC pastes [18]. A decrease in the pH conditions due to carbonation causes the decomposition of Cr(VI)-bearing hydration products in cementitious materials, which leads to more dissolved aqueous Cr(VI) [19].

Ground granulated blast-furnace slag (GGBS) is a well-known by-product of the pig-iron manufacturing process, and up to 60–70% of GGBS is often incorporated into concrete in order to recycle industrial waste materials [20]. GGBS is capable of immobilizing Cr(VI) via a reductive process: dissolved S$^{2−}$ ions from GGBS transform Cr(VI) into Cr(III) [3]. On the contrary, Macias et al. [18] observed an increased dissolution of Cr(VI) in GGBS-PC pastes due to the Cr(VI) that was released into the cement materials’ pore solution. The effect of GGBS on Cr(VI) reduction and immobilization has not yet been quantitatively studied, and the exact mechanism of GGBS remains unclear.

The aim of this study is to elucidate the mechanism of GGBS on Cr(VI) speciation under different alkaline levels. Ca(OH)$_2$ was utilized in order to maintain a high alkaline condition for the activation of GGBS. We also studied GGBS’s capacity to reduce and immobilize Cr(VI), comparing it to that of pure PC. We performed quantitative determinations of the Cr(VI) using a UV-visible spectrophotometer. The dissolved cations and anions in Cr(VI)'s reductive transformation in the solution phase were quantified under different alkaline conditions using ion chromatography. The phases for each sample that was made to react with different Cr(VI) concentrations were identified using X-ray diffraction (XRD). Synchrotron-based X-ray microscopy and spectroscopy were used to identify the detailed chemical speciation of Cr that had reacted with the Ca(OH)$_2$-activated GGBS particles. Finally, an accelerated carbonation test was performed to examine the effects of carbonation on Cr(VI) leaching from a GGBS-PC paste.
2. Experimental Methods

2.1. Material and Sample Descriptions

The chemical compositions of PC and GGBS were determined using X-ray fluorescence (XRF, Bruker, Karlsruhe, Germany) (Table 1). Research-grade PC was used instead of commercial PC to minimize any possible impurities and their subsequent interference with our experimental results.

<table>
<thead>
<tr>
<th>Materials</th>
<th>SiO$_2$</th>
<th>Al$_2$O$_3$</th>
<th>Fe$_2$O$_3$</th>
<th>CaO</th>
<th>MgO</th>
<th>SO$_3$</th>
<th>Na$_2$O</th>
<th>K$_2$O</th>
<th>TiO$_2$</th>
<th>P$_2$O$_5$</th>
<th>MnO</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC</td>
<td>20.94</td>
<td>5.45</td>
<td>2.83</td>
<td>64.96</td>
<td>1.5</td>
<td>2.05</td>
<td>0.32</td>
<td>0.48</td>
<td>0.27</td>
<td>0.31</td>
<td>0.08</td>
<td>99.22</td>
</tr>
<tr>
<td>GGBS</td>
<td>33.03</td>
<td>14.82</td>
<td>0.42</td>
<td>41.44</td>
<td>6.51</td>
<td>2.07</td>
<td>0.27</td>
<td>0.27</td>
<td>0.53</td>
<td>0.04</td>
<td>0.18</td>
<td>99.9</td>
</tr>
</tbody>
</table>

Table 1. The chemical compositions of the raw materials.

Table 2 shows the mix proportions and experimental measurements for the different samples. Four different series of experimental samples were prepared, and they are denoted as S1, S2, S3, and S4. In the S1 samples, we examined Cr(VI) reduction and immobilization using PC or GGBS. A Cr(VI) solution (0.5 mg/L) was prepared by dissolving potassium dichromate ($K_2Cr_2O_7$) in distilled water. The GGBS or PC powder (40 g) was added into the Cr(VI) solutions, and the suspensions were agitated at 20 °C in 60% relative humidity (RH) conditions. The supernatant of the S1 samples was filtered, and the concentration of the Cr(VI) remaining in the filtered solution was determined. The changes in pH of the S1 suspensions were measured via a hand-held pH meter (D-52, Horiba, Tokyo, Japan).

Table 2. The mix proportions of the specimens.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Distilled Water (L)</th>
<th>GGBS (g)</th>
<th>CH (g)</th>
<th>PC (g)</th>
<th>Cr(VI) Concentration (mg/L)</th>
<th>Measurements</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC-S1</td>
<td>2</td>
<td></td>
<td></td>
<td>40</td>
<td>0.5</td>
<td>UV-vis, XRD, pH</td>
</tr>
<tr>
<td>GGBS-S1</td>
<td>40</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GGBS-CH-S1</td>
<td>40</td>
<td></td>
<td>20</td>
<td></td>
<td></td>
<td>UV-vis, XRD, pH, Soft X-ray absorption microscopy</td>
</tr>
<tr>
<td>PC-S2</td>
<td>2.4</td>
<td></td>
<td></td>
<td>48</td>
<td>0.5</td>
<td>UV-vis</td>
</tr>
<tr>
<td>GGBS-S2</td>
<td></td>
<td>48</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GGBS-CH-S2</td>
<td>48</td>
<td></td>
<td>24</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GGBS-S3</td>
<td>2</td>
<td>40</td>
<td>20</td>
<td></td>
<td>0.5~100</td>
<td>UV-vis, XRD</td>
</tr>
<tr>
<td>GGBS-S4</td>
<td>2</td>
<td>40</td>
<td></td>
<td></td>
<td></td>
<td>Ion chromatography</td>
</tr>
<tr>
<td>GGBS-CH-S4</td>
<td>2</td>
<td>40</td>
<td>20</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

GGBS = Granulated ground blast-furnace slag; CH = Ca(OH)$_2$; PC = Portland cement; XRD = X-ray diffraction; UV-vis = Ultraviolet-visible spectrophotometry.

S2 samples were prepared in order to verify what effects the dissolved ions from PC or GGBS had on Cr(VI) reduction in the solution phase. The PC or GGBS powder (48 g) was mixed with distilled water (2.4 L), and the substrates were equilibrated in distilled water for 24 h to ensure the dissolution of the substrates. After 24 h, the supernatant was filtered, and 1.2 mg of $K_2Cr_2O_7$ was added to the filtered solutions containing the dissolved ions from the paste samples. Changes to the initial Cr(VI) concentration (0.5 mg/L) was measured at 0, 1, 3, 6, 12, and 24 h in the reacted solutions to identify the extent of the reductive reaction that was due to the dissolved ions from the substrates in the solutions.

The S3 experiments series aimed to determine the Cr(VI) sorption capacity of GGBS. Different amounts of Cr(VI) ranging from 0.5 to 100 mg/L were added to a GGBS suspension. After agitating the suspension, the remaining Cr(VI) concentration was determined at 120 and 150 h. Lastly, the S4 series was used to determine the different chemical species of the dissolved free ions from the GGBS samples under different alkaline condition. Ion chromatography ( Dionex ICS-2000, Sunnyvale, CA, USA), using suppressed conductivity detection, was performed to quantify the ions that were dissolved in S4 samples. The Cr(VI) concentration in S4 samples was measured using a UV-visible spectrophotometer (DR 5000™ , Hach,
London, ON, Canada) and the EPA 7196A method with 1,5-diphenylcarbazide (Cr(VI)) Detection limit, Max: 1.0 mg/L, Min: 0.03 mg/L. All the solutions were filtered through a 0.45-µm membrane filter, and the alkaline conditions of the solutions were maintained using pure Ca(OH)$_2$.

2.2. Accelerated Carbonation and Tank Leaching Test

GGBS-PC pastes ($\phi50 \times 100$ mm) were made at GGBS replacement ratios of 0, 5, 10, and 50 wt %. The water to total solid (GGBS + PC) ratio (w/s) of the pastes was maintained at 0.6 for all the samples. The total initial Cr(VI) concentration was at 0.5 mg/L per 100 g of paste. During the first 24 h of curing, plastic sheets covered the exposed surfaces of the specimen in the mold. The specimens were demolded after 24 h and were cured for an additional 28 days at 20 °C in 60% RH conditions. For accelerated carbonation, the final sample was finely grounded and placed in a carbonation chamber at 20 °C with 5% CO$_2$ and 60% RH for 28 days. After the carbonation period was completed, the specimens were kept in an ethanol suspension and stored in a vacuum drying chamber until further analysis. The hydration products were identified both before and after 28 days of carbonation. After carbonation, the remaining Cr(VI) concentration in the samples was measured according to Regulation No. 49 (The Ministry of Land, Infrastructure and Transport, Japan).

2.3. Phase Identification

The raw material phases, the hydration products in the filtered solids derived from the S1 and S3 suspensions, and the accelerated carbonation test specimens before and after carbonation were all identified via powder XRD using monochromatized CuK$_\alpha$ radiation ($\lambda = 1.54$ Å, at a voltage of 30 kV, a current of 10 mA, in the range of $2\theta = 5°–75°$, D2 Phaser, Bruker, Karlsruhe, Germany). The residues collected after filtering were dried and used for the XRD analysis. Prior to the sample preparation, both the carbonation of the raw materials and the purity of Ca(OH)$_2$, which was used as a reference compound, were checked using XRD. Although great care was taken to avoid carbonation of the pure Ca(OH)$_2$, very small peaks of CaCO$_3$ were seen in the XRD pattern, and the amount of CaCO$_3$ was quantified as ~2 wt % via Rietveld refinement using the Topas software (version 4.3, Bruker, Karlsruhe, Germany). This result may be due to the unavoidable carbonation of pure Ca(OH)$_2$ during the XRD measurements.

2.4. Scanning Transmission X-ray Microscopy (STXM)

Spatial correlations between the chemical compositions and their oxidation states were mapped by a combination of scanning transmission X-ray microscopy (STXM) and X-ray absorption spectroscopy (XAS) across Cr $L_{2,3}$-edges (560–600 eV) and the O K-edge (525–560 eV), which allows us to image distributions of the chemical species with spectral sensitivity from very small spots (~35 nm). STXM measurements were performed at the bending magnet beamline (5.3.2.1) at the Advanced Light Source (ALS) in the Lawrence Berkeley National Laboratory, Berkeley, CA, USA [21]. To obtain transparency in the soft X-ray region, the diluted mixtures (dried leachate of GGBS-CH-S3 and 10 mg/L of added Cr(VI)) were drop-casted with isopropanol onto a 50-nm-thick Si$_3$N$_4$ membrane (Silson Inc., Warwickshire, England). STXM measurements utilized a 25-nm outer-width Fresnel zone plate for focusing the X-ray. A raster scanned onto $8 \times 8 \mu$m$^2$ field-of-views with a square scan grid of 35 nm steps. Image spectra (i.e., repeated images at different energies), whose finest energy step was 0.1 eV near the absorption resonance, were recorded by a phosphor photomultiplier tube (PMT). The intensity transmitted through the samples ($I$) was converted into the optical density (OD) by taking the log ratio of the incident ($I_0$) and transmitted ($I$) intensities (Equation (1)):

$$\text{OD} = -\ln\left(\frac{I}{I_0}\right) = \sigma \cdot t = \mu \cdot \rho \cdot t,$$

where $I_0$: incident X-ray photon intensity, $t$: specimen thickness, $\mu$: mass absorption coefficient, and $\rho$: density of the specimen.
To overcome spectral distortions resulting from the acquisition time gap between the I₀ and I measurements, the incident beam intensity was simultaneously measured through an open area of the sample. After image registration by a phase-correlation algorithm, the XAS spectra of each pixel as a function of energy were corrected. All data processing was carried out using the Axis2000 software (version 8.3, McMaster Univ., Hamilton, ON, Canada) [22]. Further details regarding applications of STXM to the study of cementitious systems can be found elsewhere [23–26].

3. Results and Discussion

3.1. Cr(VI) Removal Using PC and GGBS-PC Samples

Figure 1a shows the time profiles of the pH values for the suspensions retrieved from the S1 samples over 120 h. The pH value for PC-S1 increased from 12.7 to 13.2 in 12 h and decreased to 12.5. This is in good agreement with PC paste, which has a water to binder ratio (W/B) of 0.3 [27]. No significant changes in pH were observed afterwards. On the other hand, the pH value for GGBS-S1 increased from 11.1 to 11.7 in 24 h and remained in the 11.6–11.8 range until 120 h had passed. When compared to PC-S1 and GGBS-S1, GGBS-CH-S1, to which Ca(OH)₂ was added, exhibited comparatively small changes in pH within the first 20 h. It had a constant value of 12.7 until 120 h had passed. Past studies indicate that GGBS’s activation is strongly dependent on pH values [28,29]. Higher pHs lead to improved GGBS activation, with a value higher than 11.5 leading to an effective activation [30]. Therefore, alkaline conditions for GGBS activation can be maintained by adding Ca(OH)₂ (GGBS-CH-S1). The effect of alkalinity on GGBS activation was further confirmed in the XRD results (see Section 3.4).

Figure 1b depicts the Cr(VI) concentration of the supernatants ([Cr(VI)]_{supernatants}) retrieved from the S1 samples as a function of time. Within 1 h, [Cr(VI)]_{supernatants} increased from 0.50 to 0.70 mg/L for PC samples. Following this, [Cr(VI)]_{supernatants} gradually decreased over 120 h to below the detection limit (0.03 mg/L). The initial increase in the Cr(VI) concentration is likely due to the initial dissolution of the cement clinkers in the PC samples. In the GGBS-S1 without Ca(OH)₂, the [Cr(VI)]_{supernatants} decreased from 0.5 to 0.4 mg/L within 5 h and remained relatively constant at 0.40 mg/L over 120 h. In contrast, a rapid decrease was observed for the GGBS-CH-S1 in Ca(OH)₂ solution, with its the [Cr(VI)]_{supernatants} reaching 0 mg/L within 24 h. This is an interesting observation because increasing the pH condition is expected to increase [Cr(VI)]_{supernatants}. Instead, we observed a [Cr(VI)]_{supernatants} decrease in the presence of the GGBS paste. Our results suggest that GGBS has an impact on Cr(VI) reaction pathways in Ca(OH)₂ conditions and that alkali-activated GGBS can remove and immobilize Cr(VI) more than the untreated GGBS in distilled water can.
3.2. Cr(VI) Reduction via Dissolved Ions from GGBS

Figure 2 shows the evolution of the Cr(VI) concentration in the filtered solutions ([Cr(VI)]\textsubscript{aqueous}) of the S2 samples as a function of time. [Cr(VI)]\textsubscript{aqueous} decreases when it reacts with dissolved ions from PC and GGBS. Water-S2, the control sample, is a mixture composed of distilled water and K\textsubscript{2}Cr\textsubscript{2}O\textsubscript{7} mixed with a pre-determined Cr(VI) concentration of 0.5 mg/L, brought to reaction. The initial [Cr(VI)]\textsubscript{aqueous} for all the samples was 0.5 mg/L. During the first hour, a slight increase in [Cr(VI)]\textsubscript{aqueous} for the PC-S2 sample (~0.52 mg/L), when compared to Water-S2 (~0.47 mg/L), was likely due to Cr(VI) dissolving from the cement clinker, which we also observed in Figure 1b because PC inherently contains a small amount of trace Cr(VI).

The initial [Cr(VI)]\textsubscript{aqueous} in GGBS-S2 decreased to 0.38 within 1 h before reaching a constant concentration. On the contrary, due to the presence of Ca(OH)\textsubscript{2}, GGBS-CH-S2 showed a significant and continuous Cr(VI) reduction over 6 h before finally reaching 0.05 mg/L at 24 h, which suggests an almost complete removal of Cr(VI) from the solution. Based on the results, we attributed the reduced amount of Cr(VI) in the solution to the solution chemistry between the dissolved ions in the GGBS-CH-S2 sample with Cr(VI). In contrast, our experimental data indicated that the dissolved ions from the PC did not reduce the Cr(VI). Our results suggested that the significant Cr(VI) concentration decrease for the GGBS in Ca(OH)\textsubscript{2} solution was primarily due to the reducing reaction between the dissolved ions from the GGBS and Cr(VI).

Figure 3 shows the evolution of the concentrations of the dissolved ions from the GGBS in water (GGBS-S4) and GGBS in Ca(OH)\textsubscript{2} solution (GGBS-CH-S4) within 25 h. In both GGBS-S4 and GGBS-CH-S4 samples, the concentrations of Al\textsuperscript{[III]} were relatively lower (lower than 1 mg/L) than those recorded for the other cations. For GGBS-S4, the three cations (Al\textsuperscript{[III]}, Si\textsuperscript{[IV]}, and Ca\textsuperscript{2+}) showed similar dissolution trends. We presumed that the concentration changes in those cations resulted from the slow dissolution of GGBS in water. The Si\textsuperscript{[IV]} concentration in GGBS-CH-S4 was very low due to the rapid consumption of Si in GGBS’s alkali-activation process.
The initial Ca\(^{2+}\) concentration in GGBS-CH-S4 (~200 mg/L) was much higher than for GGBS-S4 (~12 mg/L) because of the added Ca(OH)\(_2\). After 1 h, the Ca\(^{2+}\) concentration in GGBS-CH-S4 increased significantly to ~650 mg/L but then decreased to 400 mg/L at 3 h. After this, we recorded no changes in the Ca\(^{2+}\) concentration in GGBS-CH-S4. The initial rapid increase and the subsequent decrease of the Ca\(^{2+}\) concentration represent, respectively, the dissolution of Ca\(^{2+}\) from the GGBS and the ion’s consumption via the precipitation of alkali-activation products. The constant Ca\(^{2+}\) concentration that we recorded after 3 h resulted from the balance between the GGBS dissolution process on the one hand and the Ca\(^{2+}\) consumption via the precipitation of alkali-activation products on the other.

The \(\text{S}_2\text{O}_3^{2-}\) and \(\text{SO}_4^{2-}\) concentrations were measured to represent the anions’ dissolution form the GGBS-S4 and GGBS-CH-S4 samples. In GGBS-S4, a relatively small amount of anions were dissolved from GGBS, and this amount was constant for 25 h. Conversely, in the GGBS-CH-S4 sample, the \(\text{S}_2\text{O}_3^{2-}\) and \(\text{SO}_4^{2-}\) dissolution rapidly increased after 7 h. Most importantly, in contrast to the \(\text{S}_2\text{O}_3^{2-}\) (0–0.5 mg/L) and \(\text{SO}_4^{2-}\) (1–2 mg/L) concentrations, we observed that a significantly greater amount of total sulfur (S) (7–13 mg/L) was dissolved within 5 h (the rapid Cr(VI) removal duration for GGBS-CH-S2, in Figure 2) in GGBS-CH-S4 samples than in GGBS-S4. Previous XAS study found that the S in GGBS exists mostly in the form of \(\text{S}_2\text{O}_3^{2-}\) with a minor amount of \(\text{SO}_4^{2-}\) [31]. Our results therefore suggested that GGBS samples in Ca(OH)\(_2\) solution the significant Cr(VI)-concentration decrease in the solution phase was primarily caused by the fact that the reducing reaction between the dissolved ions (\(\text{S}_2\text{O}_3^{2-}\)) acted as a reducing agent for the Cr(VI) in the solution, as has previously been found by others [3,11]. We concluded that the dissolved ions from the GGBS activated in Ca(OH)\(_2\) solution play a critical role in the reductive transformation of Cr(VI) into Cr(III) in solution.

### 3.3. Cr(VI) Adsorption onto GGBS-PC Samples

We measured the changes in the Cr(VI) concentration for the S3 samples at 0, 120, and 150 h in order to further evaluate the capacity of GGBS to immobilize Cr(VI) in a Ca(OH)\(_2\) solution (Figure 4). For each specimen, the initial Cr(VI) concentrations varied from 0.5 to 100 mg/L. After 120 h of agitation, we observed no trace of Cr(VI) in GGBS-CH-S3 that had a Cr(VI) concentration of up to 10 mg/L, which indicated
the complete removal of Cr(VI) from that solution. The GGBS specimen containing 50 mg/L of Cr(VI) had approximately 0.7 mg/L of Cr(VI) after 120 h. The Cr(VI) concentration in the specimen containing 100 mg/L of Cr(VI) was reduced to 8.6 mg/L. Furthermore, for all samples, we observed no further change in the Cr(VI) concentrations even after 150 h. It is possible that all GGBS’s sorption sites for Cr(VI) are saturated at this concentration, which would explain why no further removal of Cr(VI) from the solution was observed. It is also possible that the saturation of GGBS’s surface sites hinders any further reduction of Cr(VI) to Cr(III). Based on there being no further changes in the Cr concentrations after over 150 h, a pseudo-equilibrium condition was attained under our experimental condition. We do not have the experimental data to measure activated complexes at the transition state for the overall Cr transformation reaction mechanisms, and as a consequence of this we cannot calculate the pseudo-equilibrium constant for the reactions. Further detailed work is needed to confirm the exact mechanism at play during surface site saturation, along with its impact on Cr(VI) uptake and reduction by GGBS. Additionally, GGBS in distilled water (GGBS-S1) exhibited a limited capacity for Cr(VI) removal within 120 h: 0.025 mg/L per gram of GGBS. Related to this result is the fact that GGBS’s capacity to immobilize and stabilize Cr(VI) in Ca(OH)2 solution was particularly large: approximately 2.3 mg/L per gram of GGBS. This result indicates that the alkaline conditions improve the dissolution of GGBS. This results in the enhancement of the GGBS’s capacity to remove Cr(VI), in addition to its potential to do so in distilled water conditions. All the reaction products of the experimental samples were further analyzed using XRD in order to examine the possible surface precipitate phase.

3.4. XRD Characterization of the Reacted Phases

Figure 5 shows the diffraction patterns of raw materials and of the S1 samples, measured using XRD. We recorded no diffraction peaks for K2Cr2O7 in S1 specimens, indicating the complete dissolution of K2Cr2O7. PC-S1 exhibited very strong crystalline peaks for the Ca(OH)2 formed from the PC hydration, but such crystalline peaks were absent in GGBS-S1’s XRD pattern, indicating that here GGBS was mostly amorphous and had not been effectively activated under a pH of circa −11.5 (Figure 1a). In the XRD patterns of PC-S1 and GGBS-CH-S1 samples, we observed a formation of calcium aluminate hemi carbonate (hereafter referred to as hemi carbonate, 2Ca4[Al(OH)6]2(CO3)3.5OH·5.5H2O) and calcium aluminate monocalc ite (hereafter referred to as monocarbonate, 3CaO·Al2O3·CaCO3·10.68H2O) while only the PC-S1 sample formed ettringite.
Based on our findings, i.e., the rapid Cr(VI) reduction and immobilization through GGBS under alkaline conditions maintained by Ca(OH)₂, we conclude that the by-products produced by the alkali-activated GGBS can contribute to both the immobilization of Cr(VI) ions via the complexation with dissolved anions resulting from the GGBS hydration reaction, and the Cr(VI) reduction to Cr(III).
3.5. Chromium Speciation via STXM

In exhibiting different Cr oxidation states, Cr(III) and Cr(VI) can be readily resolved by comparing the sample spectra with suitable reference spectra that are taken at the Cr L2,3-edge. This is because near edge X-ray absorption fine structure (NEXAFS) is highly sensitive to both the oxidation states and the geometry of Cr’s first coordination shell [32].

Figure 7 presents the X-ray absorption image of the GGBS-CH-S3 sample taken below Cr L3-edge at a photon energy of 560 eV, and the image contrast maps of Cr and O. Cr appeared to be localized at the edge of the GGBS particles, whereas O was equally distributed in the GGBS. In the Cr image contrast map, Area 1, as illustrated in Figure 8a, was selected for the Cr L2,3-edge NEXAFS analysis. We chose pure K2Cr2O7 as a reference for Cr(VI) spectra and used a series of Gaussian peaks to fit the spectra. In Figure 8b, K2Cr2O7 exhibits four distinct X-ray absorption features in the Cr L2,3-edge spectra, indicated as a2 (~578.5 eV), a3 (~580.9 eV), b2 (~587.7 eV), and b3 (~589.5 eV), where the peaks defined as a3 and b2 are induced, respectively, by the Cr L3 and L2 absorptions. The additional peaks a2 and b2 result from the crystal field produced by the ligands with different electronegativities, which are non-linearly related to the magnitude of the crystal-field parameter (10Dq) [33,34]. Since the absorption shoulder features of Cr(VI) on the low energy side are very close to Cr(III)’s main absorption [32], the absorption peaks of a2 and b2 are overlapping contributions from both Cr(VI) and Cr(III). The low energy side features a1 and b1 (the arrows in Figure 8b) solely correspond to the octahedrally coordinated Cr(III), which is in excellent agreement with the chromite (FeCr2O4) spectrum and Cr2O3 [32,35,36]. The coexistence of Cr as both Cr(VI) and Cr(III) is presumably due to the superposition of the partially-reduced Cr(VI) and the physically adsorbed Cr(VI) onto the Ca(OH)2-activated GGBS. The remaining areas illustrated in Figure 8a show the same trends as Area 1, i.e., the coexistence of Cr(III) and Cr(VI). However, due to the lower Cr concentration, the areas have lower X-ray absorption intensities and a higher background noise.

![Figure 7](image_url)

Figure 7. (a) Scanning transmission X-ray microscopy (STXM) image of GGBS-CH-S3 taken at 560 eV; (b) image contrast map of Cr; and (c) O for GGBS-CH-S3.

The O K-edge NEXAFS analysis provided further evidence for the coexistence of Cr(VI) and Cr(III) on the Ca(OH)2-activated GGBS (GGBS-CH-S3), as shown in Figure 9. The O K-edge NEXAFS spectrum of the K2Cr2O7 exhibited intense pre-edge X-ray absorption peaks at 529.2 and 531.2 eV, and these can be attributed to the electronic excitation from the O 1s core level into the unoccupied t2g and eg molecular orbitals, respectively [37]. The O K-edge NEXAFS spectrum of pure Ca(OH)2 differs from the absorption features of CrO42− [24]; in this study, the presence of crystalline Ca(OH)2 was not observed in the entire area of the specimen. Area 1 and Area 2 appeared to contain two pre-edge, low-intensity absorption peaks arising from CrO42−. In addition to the peaks, the features indicated by arrows at ~533 eV in the Area 1 and Area 2 spectra correspond to the O K-edge absorption peak of Cr(III)-oxyhydroxide, which has previously been observed in the spectrum of pyrite (FeS2) reacting...
with Cr(VI) [35]. Area 3 did not exhibit any CrO$_4^{2-}$ or Cr(III)-oxyhydroxide features because of the absence of any Cr in an oxidation state, as observed in Figure 7b.

Figure 8. (a) Illustration of the analyzed areas in GGBS-CH-S3 for the Cr $L_{2,3}$-edge NEXAFS in the Cr image contrast map (Figure 7b); (b) Cr $L_{2,3}$-edge NEXAFS spectra for Area 1 in (a) and spectra from K$_2$Cr$_2$O$_7$.

Figure 9. (a) Illustration of the analyzed areas in GGBS-CH-S3 for the O $K$-edge NEXAFS; (b) O $K$-edge NEXAFS spectra for each area in (a) and for K$_2$Cr$_2$O$_7$. 3.6. Cr(VI) Dissolution and the Effects of Accelerated Carbonation

A comparison of the dissolved Cr(VI) concentration from the specimens before and after accelerated carbonation is shown in Figure 10. At 56 days, all the specimens that had not been exposed to atmospheric CO$_2$ showed that relatively minimal amounts of Cr(VI) (i.e., less than the detection limit of 0.03 mg/L) were released from the GGBS-PC paste, regardless of the replacement...
3.6. Cr(VI) Dissolution and the Effects of Accelerated Carbonation

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For a sample with 50% of GGBS-replaced PC paste, we measure a very small amount of Cr(VI) below the detection limit of 0.03 mg/L. Pore solutions extracted from PC paste generally possess a pH in the range of 12.4 to 13.5 [38]. Our work revealed that the alkalinity resulting from the naturally formed Ca(OH)$_2$ in PC hydration was sufficient for GGBS activation and enhanced GGBS’s capacity to reduce and immobilize Cr(VI), as observed in the solution samples (Figures 1b and 2). Consequently, in a realistic PC paste, replacing PC with a minimal amount of GGBS had a significant impact on Cr(VI) reduction and immobilization, even with a 5 wt % replacement.

![Figure 10](image1.png)

**Figure 10.** Tank leaching test results for the GGBS-PC paste before and after an accelerated carbonation test.

Figure 11 shows the XRD patterns for the GGBS-PC paste, with different GGBS replacement ratios and either with or without carbonation. Peaks arising from monosulfate, Ca(OH)$_2$, and C-S-H were detected for all GGBS-PC pastes without carbonation. For the 50 wt % GGBS-PC paste, we noted very weak peaks due to the presence of hydrotalcite ($\text{Mg}_6\text{Al}_2(\text{CO}_3\text{(OH)}_{16} \cdot 4\text{H}_2\text{O})$) and vaterite. This result indicated that replacing GGBS in PC increases the hydration products’ susceptibility to carbonation due to the reaction with atmospheric CO$_2$. Previous studies have also confirmed that carbonation can change the PC paste’s porosity and that this can also affect the hardened specimen’s CO$_2$ permeability [39,40].

Monosulfate diffraction patterns were not detected in all samples with carbonation, which suggests that complete monosulfate decomposition occurred in the carbonated samples. In addition, we detected the presence of CaCO$_3$ polymorphs such as vaterite, aragonite, and calcite; these were the main, crystalline, accelerated carbonation products for all the specimens. In particular, as the carbonation progressed, the peak intensities of the CaCO$_3$ polymorphs became stronger, as has
been previously found [41]. Presumably, the formation of aragonite is linked to the carbonated C-S-H formed during PC clinker hydration, as has been observed in the young post-carbonation tricalcium silicate paste [42]. In general, Ca(OH)₂ and C-S-H are the main hydration products involved in the carbonation of PC-based systems [43]; the carbonation occurs preferentially on Ca(OH)₂, and precipitating CaCO₃ polymorphs in the system [44,45]. However, carbonation in PC systems that contain 75% amounts of GGBS mainly occurs on the C-S-H phases, which results in vaterite formation [46]. C-S-H formed in GGBS-PC systems has a generally low Ca/Si ratio because of the Ca(OH)₂ consumption compared to that of pure PC systems [47–50]. When C-S-H with a low Ca/Si ratio is exposed to CO₂, both vaterite and aragonite, which are CaCO₃ polymorphs, preferentially precipitate [51]. In the present study, after carbonation, C-S-H exhibited a peak intensity in GGBS50C that was lower (2θ = ~29°) than for those contained less GGBS. Furthermore, we observed that the more GGBS there was, the higher was the vaterite-related peak, which formed due to the carbonation of C-S-H with a low Ca/Si ratio.

Figure 11. XRD patterns of the GGBS-PC paste with GGBS replacement ratios of 0, 5, 10, and 50 wt % before and after 28 days of accelerated carbonation. ◆: Vaterite, ◇: Aragonite, ■: Ca(OH)₂, ◇: Calcite, ●: C-S-H, ★: Monosulfate, and □: Hydrotalcite.

Figure 12 shows the XRD peak intensity of the 001 basal reflection from Ca(OH)₂ for the GGBS-PC pastes before and after carbonation. We observed a very intense peak for crystalline Ca(OH)₂ in the pre-carbonation pure PC paste (GGBS0). Like anticipated, as GGBS’s replacement ratio in PC increased, Ca(OH)₂’s peak intensity decreased due to the dilution of PC and to the consumption of Ca(OH)₂ by GGBS. After carbonation, more Ca(OH)₂ decreased in the pure PC paste (GGBS0) than Ca(OH)₂ in the GGBS-PC paste. As a result, all the specimens exhibited similar Ca(OH)₂ peak intensities, which supported the findings that C-S-H is more readily carbonated than Ca(OH)₂ in GGBS-PC paste because the carbonation of C-S-H depends on the initial Ca(OH)₂ amount present prior to carbonation [46]. Moreover, it can be demonstrated that the sufficient Ca(OH)₂ in pure PC paste can buffer C-S-H carbonation. Presumably, the lower Cr(VI) dissolution in the carbonated GGBS-PC resulted from a pre-carbonation reductive process caused by dissolved anions such as S²⁻ (which are continuously released from GGBS during hydration) because S²⁻ can inhibit the oxidization of Cr(III) to Cr(VI) [18]. It should be noted that we do not have the direct evidence we need to identify the exact surface-promoted reductive transformation of Cr(VI) to Cr(III). It is likely that


S\(^{2-}\) released during the hydration reaction is responsible for the reducing reactions in solution and that the reduced Cr species are physisorbed to hydrated materials. Equally, we cannot rule out the possibility that physisorbed S\(^{2-}\) located on hydrated-materials surfaces act as reducing agents for Cr reductive transformations. Further detailed studies are needed to distinguish the two different reaction pathways. Moreover, C-S-H formed in the GGBS-substituted PC system, which has a longer mean chain length \[52\], resulting in a larger surface area that can offer better surface adsorption, inclusion, and physical entrapment of Cr(VI). Based on the XRD and Cr(VI) dissolution results before and after carbonation, we concluded that partially replacing PC with GGBS can greatly reduce and immobilize Cr(VI) via dissolved anions into the pore solutions and via physical adsorption onto C-S-H. Even though C-S-H and hydration products are more susceptible to carbonation in the GGBS-PC system than in the pure PC system, Cr(VI) was not readily leached from the GGBS-PC paste after carbonation.

![Graph showing 001 basal reflection](image)

**Figure 12.** Comparison of the 001 basal reflection from Ca(OH)\(_2\) in GGBS-PC pastes before and after an accelerated carbonation. The numbers followed by GGBS indicate the replacement ratios of GGBS for PC.

### 4. Conclusions

We found that without alkali activation GGBS is limited in its capacity to reduce Cr(VI). In contrast, the Ca(OH)\(_2\)-activated GGBS demonstrated a considerably greater capacity for reducing and immobilizing Cr(VI). Therefore, we conclude that alkalinity is one of the dominating factors in determining the extent and mechanisms of reductive Cr(VI) dissolution and uptake reactions for GGBS. Furthermore, we found experimental evidence that Cr(VI) removal from the solution was due to its reactions with the dissolved anions, such as S\(^{2-}\), resulting from GGBS being under high alkaline conditions. X-ray absorption spectroscopy data identified the coexistence of Cr(III) and Cr(VI) on the Ca(OH)\(_2\)-activated GGBS. This partial reduction of Cr(VI) to Cr(III) on the surface of GGBS suggests the presence of a surface-site-activated reductive mechanism. The Cr(VI) we observed on GGBS like results from physically adsorbed Cr(VI) on hydration products.

Our accelerated-carbonation experimental data indicate that the hydration products formed in the GGBS-PC paste, such as C-S-H, possessed a greater resistance to Cr(VI) leaching than those formed in pure PC. Our study suggests that even the used of 5% weight-based partial GGBS replacement for PC-based materials would be a suitable method for reducing and immobilizing Cr(VI) without violating the mechanical and physical integrity of PC \[53\]. We hypothesize that Ca(OH)\(_2\) generated during the early stage of PC hydration reactions promotes the dissolution and activation of GGBS, which results in an enhanced Cr(VI) reduction. It should be noted that observing S\(^{2-}\) concentrations in the dissolved solution will provide useful insights for determining and identifying further detailed
reaction pathways. Future studies that use additional information to research similar experimental conditions are necessary to identify Cr(VI)’s exact reductive transformation pathways in hydrated cementitious materials. In addition, in order to understand the Cr(VI) reductive mechanism in GGBS, further detailed mechanistic and molecular research is needed to identify the exact state of physisorbed Cr(VI) on GGBS’s hydration products, how different types and amounts of GGBS alkaline activators influence Cr(VI) reduction and immobilization, and the factors that control Cr(VI) reduction pathways in carbonated GGBS-PC materials.

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