Article

Coal Mining Waste as a Future Eco-Efficient Supplementary Cementing Material: Scientific Aspects

Moisés Frías 1,*, Rosario García 2, Raquel Vigil de la Villa 2 and Sagrario Martínez-Ramírez 3

1 Eduardo Torroja Institute for Construction Sciences (IETcc-CSIC), Madrid 28033, Spain
2 CSIC-UAM Associated Unit, Department of Geology and Geochemistry, Autonomous University of Madrid, Madrid 28049, Spain; rosario.garcia@uam.es (R.G.); raquel.vigil@uam.es (R.V.d.l.V.)
3 Institute for the Structure of Matter (IEM-CSIC), Madrid 28006, Spain; sagrario@iem.cfmac.csic.es
* Correspondence: mfrias@ietcc.csic.es; Tel.: +34-91-302-0440

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Abstract: The stockpiling of tailings around coal mines poses a major environmental problem. Nonetheless, this clay mineral (kaolinite)-based waste can be reused as a supplementary cementitious material (recycled metakaolinite) in the manufacture of future eco-efficient cements. This paper explores the most significant scientific questions posed in connection with the conversion of this waste into pozzolans, such as the variation in product mineralogy depending on the sintering temperature and its effect on reaction kinetics in the pozzolan/Ca(OH)2 system over a period of 365 days. The findings show that the optimal sintering temperature is 600 °C, such that the cementitious properties of the activated product are determined solely by the conversion of kaolinite into metakaolinite and are unaffected by the other clay minerals (micas). The presence of 20% activated coal waste favors the formation of larger amounts of aluminous phases such as C4AH13 and C4A3H12 than in the reference paste and enhances C–S–H gel polymerization.

Keywords: coal mining waste; mineralogy; blended cements; hydrated phases

1. Introduction

Today’s society is undergoing a change from a linear (produce-use-discard) to a recycling-based circular economic model that pursues two ends: the generation of less waste and its reuse as raw materials in industrial processes. This sustainable approach is one of the mainstays of Europe’s 2020 strategy [1].

Cement manufacturing is a benchmark in this regard, in light of its use decades-old use of industrial by-products and waste in different stages of production: as alternative fuels (worn tires), raw materials (waste water treatment sludge) or supplementary cementitious materials (SCMs) to produce eco-efficient cements [2–4]. Metakaolinite, a material well known for its high pozzolanicity that can be obtained by thermally activating natural kaolinite, is one of the standardized SCMs [5–11]. Products with a high metakaolinite content can also be obtained by activating high-kaolinite industrial waste, a more eco-friendly approach compliant with the fundamental premises of the circular economy [12–14].

The stockpiling of kaolinite-based coal mining waste (CW) has been posing severe environmental problems for decades [15–17]. These tailings are one of the primary sources of pollution in China [18], where around 4.5 billion tons have accumulated in over 1700 dumps, together occupying an area of 150 km2. In light of its kaolinitic nature, if recycled, this carbon waste may be an alternative source of MK [19–21]. Prior research conducted by Frías et al. [22,23], García et al. [24] and Vegas et al. [25] has
shown that this inert waste can be converted by thermal activation into a high added value pozzolan (activated coal waste, ACW) apt for use in the manufacture of innovative eco-efficient cements. The need has thus arisen to determine the short- and long-term behavior of these pozzolanic materials in blended cements.

This study expands on the existing scientific knowledge of both preliminary thermal activation and the subsequent hydration of these materials when blended with cement. A number of instrumental techniques (XRD, SEM-EDX, TG/DTA, micro-Raman and NMR) are deployed to analyze the phases initially present in the thermally activated MK and in 365 days hydrated blends containing 80% cement. The findings are compared to the results for a reference Ordinary Portland Cement (OPC) paste to determine the effect of the pozzolanic reaction between the ACW and the portlandite in the blended cement system.

2. Experimental Section

2.1. Materials

Raw coal mining waste (CW) was collected in the vicinity of an open-pit mine owned by Sociedad Anónima Hullera Vasco-Leonesa and located in the Spanish province of Leon. The waste was crushed, sieved under 90 microns and thermally activated at temperatures ranging from 600 °C to 900 °C for 2 h to produce ACW. Laser granulometry results showed that 10%, 50% and 90% of the ACW particles are less than 1.7, 8.11 and 32.63 microns, respectively.

The cement used was a commercial ordinary portland cement classified as CEM I 52.5 N [2]. The potential composition of the Portland clinker, computed as stipulated in the existing Spanish legislation [26], is given in Table 1. The blended cements contained 20% ACW, the maximum replacement ratio allowed for the manufacture of type II/A cements (6%-20%). After removal from the molds, the 1 cm × 1 cm × 6 cm prismatic specimens were cured in water at 20 °C until tested at 1, 28 or 365 days.

Table 1. Potential composition of clinker.

<table>
<thead>
<tr>
<th>(%)</th>
<th>C₃S</th>
<th>C₂S</th>
<th>C₃A</th>
<th>C₄AF</th>
</tr>
</thead>
<tbody>
<tr>
<td>OPC</td>
<td>43.61</td>
<td>27.71</td>
<td>9.29</td>
<td>9.70</td>
</tr>
</tbody>
</table>

2.2. Analytical Methods

The accelerated chemical method (40 °C) was applied to assess the pozzolanicity of the carbon waste activated for 2 h at 600 °C in an electrical muffle furnace. Further to that method, 1 g of pozzolan was immersed in 75 mL of saturated lime solution for a given reaction time. The solution was subsequently filtered and the CaO concentration was calculated in both the problem and control solutions. The sole difference between this method and the standardized pozzolanicity test for pozzolanic cements lies in the material, which was a pozzolan rather than a cement.

2.3. Instrumental Techniques

The mineralogical composition of the bulk samples was determined by X-ray powder diffraction (XRD) and the <2 μm fraction by the oriented film method, both on a Siemens D-5000 (Munich, Germany) X-ray diffractometer fitted with a Cu anode. Semi-quantitative mineralogical composition was calculated with the powder reflection method. The areas under the reflections for each mineral were found with Gaussian fitting, in which conventional software was applied to subtract the baseline intensity. The relative error for this method was calculated at 10%.

Sample morphology and microanalysis were conducted on an FEI Inspect (Hillsboro, OR, USA) scanning electron microscope equipped with a tungsten (W) source DX4i dispersive X-ray analyzer and an Si/Li detector. The chemical composition was determined as the mean of ten scans per sample,
shown along with the standard deviation. The chemical formulas for the minerals were determined with EDS semi-quantitative analysis and the results expressed in oxides (wt.%).

TG/DTA trials were run on a Stanton Equipment Inc. STA 781 thermogravimetric analyzer. Powder samples weighing 30 mg to 35 mg were heated at a rate of 10 °C/min in an N₂ atmosphere.

Dispersive Raman spectra were recorded at 532 nm (Nd:YAG) on a Renishaw InVia system fitted with a Leica confocal Raman microscope and an electrically cooled CCD camera. Frequency was calibrated to silicon (520 cm⁻¹) and the spectral resolution was set at 4 cm⁻¹.

The Bruker AVANCE-400 (9.4 T) spectrometer used for 29Si NMR analysis operated at a 29Si frequency of 79.4 MHz. Kaolin (d = −91.5 ppm) referenced to TMS (d = 0 ppm) was used as the external standard for chemical shift. The delay time was 60 s and the spinning speed 4 kHz.

3. Results and Discussion

3.1. Comparison of Mineralogy after Activation at 500 °C to 900 °C

According to the XRD findings, the CW sample contained phyllosilicates (micas and kaolinite), quartz, calcite, dolomite and feldspars (Figure 1). Pozzolanic activity was induced by the phyllosilicates, for thermal activation in air at 600 °C to 900 °C is known to prompt dehydroxylation in many clay minerals [27,28], with the concomitant total or partial breakdown of their crystalline lattice structure and the formation of a highly reactive transition phase.

![Figure 1](image-url) X-ray powder diffraction (XRD)-determined mineralogical composition (%) for the starting (CW) and thermally treated (ACW 500/900) coal waste.

Clay is most reactive when the activation-induced loss of hydroxyl ions leads to structural collapse and disarrangement. The sintering temperature required to convert inert kaolinite into activated metakaolinite (MK) usually ranges from 600 °C to 800 °C. Mica, in contrast, must be heated to over 930 °C to be activated and even then yields weak pozzolans.

The waste was heated at 500 °C, 600 °C, 700 °C, 800 °C and 900 °C to determine the optimal temperature for phyllosilicate dehydroxylation and new phase formation. The XRD results showed that full kaolinite dehydroxylation took place at 600 °C, with spinel-like phase neoformation occurring at 900 °C (Figure 1). The metakaolinite clusters visible in the SEM micrographs of ACW-600 were spongy and had blurred edges, whereas in the ACW-900 product the MK clusters were compact and had very sharply defined edges (Figure 2).
Consequently, 600 °C was the temperature required to convert the kaolinite in coal waste into metakaolinite.

3.2. Pozzolanic Activity

The amount of lime fixed by the activated waste over time (found with the accelerated chemical method described in the experimental section) was expressed as the ratio between the CaO content (in mmol/L) in the problem solution and the content in the control solution to obtain the pozzolanic index (PI) at each test age. According to the data in Table 2, waste ACW-600 exhibited high pozzolanicity, because 80% of the lime had been consumed in the first 28 days. The reaction rate declined after that age, with a PI of only 89% after 365 days. The behavior of ACW-600 was comparable (except in the first 24 h) to that of other types of industrial waste such as silica fume (SF) and bamboo leaf ash (BLA), regarded in the literature to be highly reactive pozzolans [29–31].

Table 2. Pozzolanic index (%) for ACW-600, bamboo leaf ash (BLA) and silica fume (SF) by hydration time.

<table>
<thead>
<tr>
<th>PI (%)</th>
<th>ACW-600</th>
<th>BLA</th>
<th>SF</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 day</td>
<td>44.6</td>
<td>81.9</td>
<td>83.1</td>
</tr>
<tr>
<td>28 days</td>
<td>80.3</td>
<td>90.9</td>
<td>88.2</td>
</tr>
<tr>
<td>90 days</td>
<td>86.7</td>
<td>91.4</td>
<td>88.2</td>
</tr>
<tr>
<td>365 days</td>
<td>89.9</td>
<td>92.1</td>
<td>90.1</td>
</tr>
</tbody>
</table>

3.3. Hydrated Phase Behavior

3.3.1. XRD Analyses

XRD analysis detected portlandite, ettringite, tetracalcium aluminate hydrate (C₄AH₁₃) and tetracalcium aluminate carbonate hydrate (C₄AcH₁₂) in both the OPC and the 20% ACW blended cement (Figures 3 and 4, respectively). The proportions of each crystalline hydrated phase varied with hydration time and the presence or absence of the ACW.

Portlandite was the predominant hydrated phase in OPC at all test ages, followed in the first 28 days by ettringite. Tetracalcium aluminate carbonate hydrate (C₄AcH₁₂) formation was clearly visible after the first 28 days and the content of this phase remained flat throughout the period studied. Tetracalcium aluminate hydrate (C₄AH₁₃) was detected at 365 days. Portlandite declined substantially after 365 days and ettringite only slightly, while calcite grew.
As in OPC, portlandite was the majority phase in the 1 day 20% ACW. Quartz, absent in OPC, was detected in the blended cement. The pattern of variation with reaction time of ettringite, tetracalcium aluminate carbonate hydrate (C₄AcH₄₂) and tetracalcium aluminate hydrate (C₄AH₁₃) also differed in 20% AWC relative to OPC.

Ettringite was the second-most abundant phase in the latter paste only up to day 1, when C₄AcH₄₂ and C₄AH₁₃ formation was clearly visible. The inclusion of 20% ACW favored the formation of C₄AH₁₃, which was the majority phase at 28 days. Its appearance was attributed to the supersaturation of the aqueous phase with calcium hydroxide. The high concentrations of Ca²⁺ and OH⁻ in the solution due to the presence of Ca(OH)₂ favored the precipitation of C₄AH₁₃, although the content of this phase declined in the 365 days samples.

Calcium moncarboaluminate is normally detected immediately after hydration. The reaction between C₃A and CaCO₃ is governed by a solid-state mechanism, while the presence of the latter modifies the initial swift hydration of the former as a result of the barrier quickly generated by the reaction product, calcium carboaluminate hydrate. As hydration progresses, hemicarboaluminate gradually converts into moncarboaluminate. That process was not observed in the present study, however, because the first analysis was conducted 24 h after the materials were mixed with water. Calcite content grew with time.

3.3.2. SEM/EDX Analyses

The combined use of these two techniques afforded information on the morphology, texture and chemical composition of the hydrated phases, and particularly about the phases that could not be identified with XRD. SEM identified C–S–H gels (Figure 5). In the first 365 days, their CaO/SiO₂ ratio ranged from 1.6 to 2.5, therefore constituting Taylor classification type II gels. The ratio was closer to 2.5 in the OPC gels and lower in the gels present in the blended cement. Structural order
rose with higher Ca/Si moduli and longer reaction times. In the early age pastes (1 day), the C–S–H gels comprised flaky clusters (Figure 5B), whereas at later ages (28 days) they exhibited thin layers of C\textsubscript{4}A\textsubscript{2}H\textsubscript{12}, C\textsubscript{4}AH\textsubscript{13} and portlandite intergrowths (Figure 5A). All three products were consistently laminar although their sizes varied. At longer hydration times (365 days), thin layers of portlandite aggregated into more compact clusters (Figure 5D), while ettringite prisms grew in the voids, together with C–S–H gel and thin layers of C\textsubscript{4}A\textsubscript{2}H\textsubscript{12} and C\textsubscript{4}AH\textsubscript{13}.

![Figure 5](image-url)

**Figure 5.** SEM micrographs of OPC and 20% ACW cement pastes at different curing times: (A) portlandite, (B) CSH gel, (C) calcite and (D) portlandite.

The inclusion of 20% ACW induced a more compact microstructure in the blended cement cured at 365 days, which exhibited larger and thicker layers of C\textsubscript{4}AH\textsubscript{13} and portlandite. Moreover, the C–S–H gel and prismatic ettringite clusters were less porous in the blended paste, in which rhombohedral calcite crystal content was likewise significant (Figure 5C).

### 3.3.3. TG/DTA

According to the thermal differential analyses (Figure 6), both cements exhibited three main endothermal signals. The wide band at 20 °C to 200 °C was attributed to water loss (45–50 °C); the band at 95 °C to 100 °C to ettringite and C–S–H gel dehydroxylation; and the band at 130 °C to 200 °C to the dehydroxylation of silicoaluminate (C\textsubscript{2}ASH\textsubscript{8}) and tetracalcium aluminate (C\textsubscript{4}AH\textsubscript{13}) hydrates, characteristically found in metakaolinite-based pozzolans [32,33]. Further to the XRD and SEM findings, the peak at 140 °C can be attributed primarily to C\textsubscript{4}AH\textsubscript{13} formation. The content of this hydrated phase rose with reaction time, particularly where carbon waste was present in the cement.
paste. No losses were identified at temperatures of around 200 °C, which would be attributed to carbo-aluminate phase (C₄AcH₁₂) decomposition.

Figure 6. DTA curves for OPC and 20% ACW by hydration time.

Portlandite dehydroxylation was observed at 400 °C to 500 °C. The Ca(OH)₂ contents from TG curves and the portlandite content (fixed lime: 100 – (20% ACW/OPC × 100)) in the pastes containing 20% ACW compared to OPC is given in Table 3 for the three reaction times studied.

Table 3. Ca(OH)₂ contents (%) and difference in both cements.

<table>
<thead>
<tr>
<th>Cements</th>
<th>1 Day</th>
<th>28 Days</th>
<th>365 Days</th>
</tr>
</thead>
<tbody>
<tr>
<td>OPC</td>
<td>13.7</td>
<td>17.9</td>
<td>19.4</td>
</tr>
<tr>
<td>20% ACW</td>
<td>12.5</td>
<td>15.7</td>
<td>11.3</td>
</tr>
<tr>
<td>Difference</td>
<td>-8.5</td>
<td>-12.1</td>
<td>-41.8</td>
</tr>
</tbody>
</table>

Pursuant to these data, the ACW clearly exhibited pozzolanic activity after 28 days due to the pozzolanic reaction between the metakaolinite and the portlandite generated during portland cement hydration, with the formation, primarily, of additional amounts of C–S–H gels and C₄AH₁₃, as discussed earlier.

The endothermal band at 650 °C to 700 °C was attributed to decarboxylation of the calcite present in both the starting OPC and the partially carbonated samples (an effect of curing and storage).

3.3.4. Micro-Raman

The 365 days Raman spectra for the OPC and 20% ACW pastes are shown in Figure 7. Both exhibited signals generated by characteristic cement hydration products: C–S–H gel (668 cm⁻¹), ettringite and monosulfoaluminate (999 cm⁻¹ to 980 cm⁻¹). Whereas the signal peaking at 1084 cm⁻¹ and assigned to calcite was narrow on the OPC spectrum, in the spectrum for the sample containing 20% waste it was wider (1070–1050 cm⁻¹) due to the presence of carboaluminate-like compounds. The OPC paste spectrum also contained a signal attributable to the OH groups in unreacted portlandite.

The ²⁹Si NMR spectra for the 365 days samples recorded to determine the characteristics of the C–S–H gel formed are reproduced in Figure 8. The weak signal observed at −71 ppm indicated that, despite the time elapsing, both pastes contained anhydrous phases. The main difference between the two spectra was that the Q₁ unit signal was much more intense in the OPC than in the 20% ACW cement paste, denoting a smaller proportion of end of chain units and hence a higher degree of polymerization in the latter. Other authors have reported similar results for the pozzolanic reaction in fired clay-based waste [34]. Furthermore, the amount, distribution and microstructural characteristics of the C–S–H gel formed have a significant effect on the mechanical strength of the cement. Studies conducted by Tobón et al. [35] showed that strength rises with the amount of C–S–H gel. The presence of over 5% pozzolanic materials (which generate more polymerized gels), in turn, raised blended cement compressive strength substantially.
4. Conclusions

As a result of its kaolinitic nature, coal mining waste affords an alternative avenue for obtaining metakaolinite, with substantial environmental and social benefits, one of the main pillars of the European circular economy.

The optimal sintering temperature for thermally activating coal waste, 600 °C, induces phyllosilicate dehydroxylation, giving rise to pozzolanic metakaolinite. When the temperature reaches 900 °C, in contrast, spinel-like phases form, lowering pozzolanicity. The amorphism and porosity of the metakaolinite clusters formed afford the material a high pozzolanic index, comparable to the indices for other highly reactive pozzolans.

Further to the mineralogical, chemical and microstructural findings discussed above, ACW is apt for use as a pozzolanic material in blended cements. The addition of 20% ACW to OPC intensifies C₄AcH₁₂ and C₄AH₁₃ formation. In the presence of the activated waste, C₄AH₁₃ is the majority reaction product. Its higher content induces more compact microstructures with a greater abundance of over 5% pozzolanic materials (which generate more polymeri...
of thin layers of C₄AH₁₃ and portlandite and lesser amounts of porous C–S–H gel and ettringite clusters. The C–S–H gel forming in OPC may exhibit fairly high Ca/Si ratios.

Economically, energetically and mineralogically speaking, coal tailings are apt for manufacturing eco-efficient cements, for at 600 °C kaolinite is fully converted to metakaolinite, whose hydration generates the same amorphous and crystalline compounds as ordinary cement.

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

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