Conference Paper

Preliminary Study on the Influence of Different Carbonation Curing Processes on Binders Based on Magnesium Oxide-Rich Powder Blended with Tungsten Mining Waste Mud

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Abstract

In this preliminary study, the effect of the pre-drying stage, water immersion, carbonation curing cycles, and/or drying stage on carbonation curing of magnesium oxide-rich powder (MRP) was investigated. In addition, a blend of tungsten mining waste mud (TMWM) with MRP was also evaluated. The MRP and TMWM used have maximum grain sizes of 125 μm. The cement pastes were produced with 0 and 50 of TMWM weight percentage. The specimens were compacted into cylindric moulds (Ø = 20 mm; h = 40 mm) under 30 MPa and, subsequently, submitted to five different processes of curing involving a pre-drying stage before carbonation, rapid water immersion cycles, additional drying periods, and different carbonation curing periods. The atmosphere of the pressurized carbonation curing chamber was controlled to provide a CO₂ concentration of > 99%, the partial pressure of 1 bar and temperature of 60°C. The influence of the curing processes on the compressive strength of each mix was determined 12 hours after the carbonation curing period. This study demonstrates that the water content during the curing process plays an important role in the increase of the hardening process and on the compressive strength.

Keywords: Carbonation curing, magnesium oxide, mining waste, curing processes, magnesium-based cement

1. Introduction

The level of anthropogenic greenhouse gas emissions is higher than ever. This level has increased since the pre-industrial era, mainly due to the economic and population growth and the last four decades comprise half of the anthropogenic CO₂ emitted between 1750 and 2011 [1]. In the midst of this, the cement industry is responsible for approximately 7% of the CO₂ emissions worldwide [2]. Therewith, the need to mitigate the CO₂ emissions linked to the conventional cement production allied to the concerns...
on the climate changes has given some popularity to magnesium-based cements, mainly over the last decade [3]. In this scope, one of the main types of magnesium-based cement is the reactive magnesia cement (RMC), which the literature describes as a blend of reactive MgO powder with conventional PC systems [3]. Such cement class is based on the carbonation cure for compressive strength gain, where the magnesium oxide (MgO) hydrates forming brucite, which is subsequently carbonated turning into hydrated magnesium carbonates (HCM), such as nesquehonite, hydromagnesite, and dypingite [4]. Some MgO-based cements have many properties that are superior to those of ordinary Portland cement (OPC), such as: good resistance to fire; high early strength; match very well to a wide variety of inorganic and organic aggregates, like gravel, marble flour, sand, sawdust [5], asbestos, wood particles and expanded clays [6]. Allied to it, there is the advantage of CO$_2$ activated binders, which have the potential to transform waste into building construction material, to store CO$_2$ permanently and also stop these resources from ending in landfills [7].

It is important to know the factors that control the carbonation reactions and also understand how it affects its kinetics. Such factors have a great influence on its efficiency and two of them are temperature and partial pressure since the atmosphere's temperature influences the CO$_2$ sequestration level, which enhances until the temperature reaches 60°C [8], and a high atmosphere partial pressure allows the CO$_2$ to enter more easily into the cementitious interior matrix before occurring the precipitation of carbonates, thus contributing to the improvement of CO$_2$ diffusivity in the cementitious matrix. [9–11].

Therefore, in this preliminary study, the influence of the water content on the carbonation curing was investigated and a method of pre-drying treatment and water immersion cycles were proposed aiming to improve the carbonation curing efficiency on magnesium oxide-rich powder (MRP) as well on a blend of tungsten mining waste mud (TMWM) with MRP, to access their effect on the hardening process and the compressive strength. For this purpose, five different curing processes were executed on two different mixes (one with 100% MRP and other with 50% MRP and 50% TMWM). The use of TMWM is supported since this waste comes from Panasqueira mines, an important mine located in the Beira Interior region, Portugal, which has been operating for over 125 years. It produces huge amounts of waste daily which come mainly from mining tungsten. Such wastes have the potential to be considered a new raw material since the industrial applications are not limited by the cost of its production and/or transportation [12]. In addition, some studies demonstrate that TMWM from Panasqueira mines can be used as raw material on cementitious blends with other different technologies [13, 14].
2. Materials and Methods

2.1. Materials

The magnesium oxide-rich powder (MRP) used is a calcinated magnesia, whose commercial name is “Magal P”, supplied by Invivo nSA, which is mainly used as an animal feed additive. Before being used, the received material was grinded in a ball mill grinder and sieved to obtain particles of a diameter size under 125 μm.

The Tungsten Mining Waste Mud (TMWM) was obtained by collecting it from the Panasqueira mine mud waste deposits. Before being used, the TMWM was disaggregated and sieved in order to obtain particles of diameter sizes under 125 mm.

To determine the chemical compositions of MRP and TMWM, Energy-dispersive X-ray spectroscopy (SEM-EDX) tests on a Hitachi S-3400N apparatus were conducted, where three different spots were randomly selected on each material sample to obtain their chemical compositions, which are presented in Table 1.

The Loss on ignition (LOI) of MRP and TMWM were obtained by simultaneous thermogravimetric and differential scanning calorimetry (TG-DSC) using a TA Instrument SDT Q-50 apparatus. The MRP lost approximately 1.82% of its weight, while the TMWM lost about 11.34% of its weight at the temperature of 1000 °C as it is shown in Table 1.

To obtain the Blaine fineness of MRP and TMWM, three tests in each sample of the raw material were conducted in an Acmel Labo BSA1 apparatus according to BS EN 196-6 [15], and then the average value was calculated. The Blaine specific area of MRP (918 cm²/g) is slightly higher than TMWM (850 cm²/g).

The particle size analyses were carried out in accordance with BS 1377-2:1990 by sedimentation using the hydrometer method [16]. The particle size distribution curve is presented in Fig. 1. The D50 of MRP is equal to 14.5 μm and D50 of TMWM is 19.5 μm.

2.2. Mix design

The mix compositions of all prepared pastes are detailed in Table 2. The weight percentage (wt. %) of MRP varies from 100% to 50% while the TMWM varies from 0% to 50%. By contrast, the water to solid (w/s) ratio was kept constant in 0.25.
TABLE 1: Chemical composition (%) and physical property of raw materials.

| Chemical composition (%) and physical property | Material | MRP | TMWM |
|-----------------------------------------------|---------|-----|------|
| SiO₂                                          |         | 6.36| 46.67|
| Al₂O₃                                         |         | 1.38| 17.05|
| Fe₂O₃                                         |         | 3.12| 15.47|
| CaO                                           |         | 3.36| 0.69 |
| MgO                                           |         | 85.78| 4.83 |
| K₂O                                           |         | -   | 4.9  |
| Na₂O                                          |         | -   | 0.8  |
| SO₃                                           |         | -   | 79   |
| TiO₂                                          |         | -   | 0.6  |
| ZnO                                           |         | -   | 1.09 |
| Loss on ignition (%)                          |         | 1.82| 11.34|
| Specific gravity (g/cm³)                      |         | 3.45| 3.19 |
| Specific surface area (cm²/g)                 |         | 918 | 850  |
| D₅₀ (mm)                                      |         | 0.0145| 0.0195|

Figure 1: Particle size analyses of raw materials
2.3. Specimens preparation

The MRP and TMWM were mixed until it resulted in a homogeneous powder and then the liquid was gradually added into the mixture until a uniform paste was achieved. Later, the prepared paste was compacted into cylindric moulds (Ø=20 mm; h=40 mm) using 30 MPa of static compaction pressure. During the static compaction stage, part of the water present in the mix is expelled, thus reducing significantly the water content on the moulded specimens. Finally, the moulded specimens were subjected to five different CO₂ curing processes.

2.4. Sample's weight and water content

The water content was determined before starting the curing processes. The water content of the fresh moulded specimens produced was determined using an electronic moisture meter KERN MLB_N apparatus, while the weight measurement of the moulded specimens was carried out by a precision balance readability of 0.01g, which has been taken the medium value of four specimens.

2.5. Pre-drying stage preliminary study

To understand the influence of a pre-drying stage (PDS) on the hardening process, several moulded specimens of TMWM00C were submitted into an oven for a PDS, during different drying periods (0, 15, 30, 45 and 60 minutes) at 60°C and afterwards submitted to carbonation curing for 2 hours at controlled atmosphere (CO₂ concentration > 99%; partial pressure = 1 bar; temperature = 60°C). Thereafter, those specimens were broken longitudinally, the core was analysed, as well as the specimen's water content. Among these, the specimens submitted to 45 min of pre-drying stage exhibited an uniformly distributed and best-hardened core, showing a less lose powder quantity inside the core. In these specimens, the water content reduced from 13.60% (before PDS) to 10.9%.
after 45 min of PDS (Table 3). Thus, based on this preliminary study a PDS of 45 min was adopted in this study.

| Time (min) | Water content | Initial | Final |
|-----------|---------------|---------|-------|
| 0         | 13.60%        | 13.6%   |
| 15        | 12.6%         |         |
| 30        | 11.2%         |         |
| 45        | 10.9%         |         |
| 60        | 8.2%          |         |

2.6. Curing processes

Five different curing processes were adopted. On the curing process 1 (CP1), the samples, after being moulded, were subjected to a carbonation curing period (CC) of 6h. The curing process 2 (CP2) involves a PDS for 45 min followed by CC of 6h. The curing process 3 (CP3) comprises a PDS for 45 min and a CC for 2h, followed by one cycle of water immersion (WI) for 5 seconds and CC for 4h. The curing process 4 (CP4) introduces the second cycle of WI and CC, which embraces a PDS for 45 min and a CC for 2h, followed by two subsequent cycles of WI for 5 seconds and CC for 2h. The curing process 5 (CP5) introduces the drying stage (DS) after water immersion, which embraces a PDS for 45 min and a CC for 2h, followed by two subsequent cycles of WI for 5 seconds and CC for 2h. However, if the specimen's weight after WI is higher than the specimen's weight after PDS, then the specimen is submitted to DS to lose the extra water content obtained on the water immersion cycle. Every carbonation curing processes was done at controlled atmosphere (CO₂ concentration > 99%; partial pressure = 1 bar; temperature = 60°C).

Thereafter, the specimens were preserved on room conditions (temperature 20 ± 2 °C; relative humidity ± 60%) for 12 hours until being tested. The control specimens (TMWM00C) were prepared to provide a comparison with specimens that have 50% of TMWM on its weight composition.
### Table 4: Carbonation curing processes

| Curing process | PDS | CC | 1st cycle | 2nd cycle |
|---------------|-----|----|-----------|-----------|
|               |     |    | WI | CC | WI | CC |
| CP1           | -   | 6 h | -  | -  | -  | -  |
| CP2           | 45 min | 6 h | -  | -  | -  | -  |
| CP3           | 45 min | 2 h | 5 s| 4 h| -  | -  |
| CP4           | 45 min | 2 h | 5 s| 2 h| 5 s| 2 h|
| CP5           | 45 min | 2 h | 5 s*| 2 h| 5 s*| 2 h|

* Followed by drying stage (DS)

**Figure 2**: Specimens of TMWM00C and TMWM50C soon after the CP5.

#### 2.7. Compressive strength

The compressive strength tests occurred 12h after the carbonation curing of 6 hours, which were measured by uniaxial loading in triplicates on the two mix compositions of the five curing processes. The equipment used was an ADR Touch 3000 BS EN Compression Machine with Digital Readout and Self Centring Platens, operated at a constant loading rate of 0.5 kN/sec.

#### 3. Results and Discussion
3.1. Water content

The TMWM00C presented 11.5% as initial water content, while the TMWM50C presented 13.9%. The PDS decreased the water content by 1% on TMWM00C and 0.8% on TMWM50C.

It was noticed that during the carbonation curing of TMWM00C only the CC of the $2^{nd}$ cycle on the CP5 did not present loss of water content. So, it is assumed that the atmosphere medium could be saturated which in turn has inhibited the loss of water from the specimen TMWM00C-CP5 on the last curing cycle. The specimen TMWM50C-CP5 also presents a similar behaviour, but in this case, it has presented a gain of water content on the CC of the $1^{st}$ and on the $2^{nd}$ cycle on the CP5.

Every specimen has shown loss of water content 12h after the curing process has finished. Such behaviour indicates that the specimens still hold excess free water on themselves after carbonation curing.

The highest loss of water content after carbonation curing that happens on TMWM00C occurred in CP3, while on TMWM50C occurred in CP2.

The results are summarized in Table 5 and Table 6, and in Fig. 3 and Fig. 4.

| Curing process | CP1 | CP2 | CP3 | CP4 | CP5 |
|----------------|-----|-----|-----|-----|-----|
| Initial        |     |     |     |     |     |
| PDS            | -   | 10.5| 10.6| 10.6| 10.4|
| CC             | 9.8 | 9.7 | 9.9 | 9.8 | 9.7 |
| Cycle          |     |     |     |     |     |
| $1^{st}$ WI    | -   | -   | 10.8| 10.6| 10.8|
| DS             | -   | -   | -   | -   | 10.3|
| CC             | -   | -   | 9.6 | 9.7 | 9.6 |
| $2^{nd}$ WI    | -   | -   | -   | 10.6| 10.4|
| DS             | -   | -   | -   | -   | 10.3|
| CC             | -   | -   | -   | 10.0| 10.5|
| 12h after curing | 9.4 | 9.5 | 9.3 | 9.5 | 10.3 |

3.2. Compressive strength

The specimens cured by carbonation curing achieved compressive strengths up to 29.7 MPa, where the best compressive strength belongs to the specimen TMWM00C-CP2. The compressive strength of the specimens that were submitted to CP3 and CP4 was
Table 6: TMWM50C water content by curing process (%)

| Curing process | CP1 | CP2 | CP3 | CP4 | CP5 |
|----------------|-----|-----|-----|-----|-----|
| Initial        | 13.9|     |     |     |     |
| PDS            |     | 13.2| 13.1| 12.9| 13.1|
| CC             | 13.4| 12.8| 11.9| 12.1| 11.9|
| Cycle WI       |     |     | 14.0| 14.0| 14.1|
| DS             |     |     |     | 12.9|     |
| CC             |     |     | 13.3| 13.4| 13.5|
| Cycle 2nd WI   |     |     |     | 14.5| 14.2|
| DS             |     |     |     | 13.4|     |
| CC             |     |     |     | 13.4| 14.0|
| 12h after curing | 13.0| 12.7| 12.8| 12.7| 13.5|

Figure 3: TMWM00C water content along the time (%)

lower than the minimum value that could be read. On the other hand, the specimens TMWM00C and TMWM50C of CP1, CP2, and CP5 compressive strength are presented in Fig. 5, which shows the comparison of compressive strength of the composition of different blends versus the curing process which was submitted.

The specimen TMWM00C suggests that the MRP reacts with the water present in the mix and also with the CO₂. Such behaviour was already expected since the MgO present on MRP reacts with water forming magnesium hydroxide (Mg(OH)₂), which reacts with dissolved CO₂ on the water forming hydrated magnesium carbonates (HMCs) [4].
The PDS shows that this stage improves the compressive strength of the proposed mix labels since both presented a strength gain after PDS.

The excess of water content does not collaborate with the gain of strength since the specimens which received the WI step and were not followed by the DS (CP3 and CP4), presented compressive strength lower than the equipment can read. On the other hand, if the excess of water is controlled, as it was done in CP5, the compressive strength increases if compared with CP3 and CP4.

The compressive strength of CP1 and CP5 has not presented a significant difference.

The mix designs have not presented a significant effect on the compressive strength (reduction less than 15% for 50% wt. % of TMWM), therefore it may assume that the carbonation conditions have much more influence on the compressive strength of the binders produced than the addition of 50 wt. % of TMWM. Thus, it may suppose that part of the MgO present in the mixes has not reacted.

4. Conclusions and Future Studies

This preliminary study has evaluated the carbonation reaction and hardening of a series of specimens of magnesium oxide-rich powder (MRP) and tungsten mining waste mud (TMWM) at different moisture conditions during carbonation curing. The water content was analysed, as well as the compressive strength of 100% MRP and 50% MRP + 50%
TMWM blends produced on five different curing processes. Therefore, the results of this study are summarized below:

1. Initial water content, as well as the water content during the curing process, plays an important role in carbonation curing and consequently on compressive strength.

2. The carbonation processes conditions had more influence on the compressive strength than the addition of 50 wt. % of TMWM, since the compressive strength difference between the mix designs are lower than 15%. Thus it suggests that part of the MgO present in the mixes had not reacted.

3. The excess of not controlled water content is inadequate to carbonation curing since the specimens submitted to CP3 and CP4 demonstrated it.

4. Initial PDS reveals itself beneficial for water content control since CP2 presented higher compressive strength than CP1 and the only difference between these curing processes is that the PDS is present on CP2;

5. The DS executed on CP5 also reveals itself beneficial for water content control since the only difference between the CP4 and CP5 curing processes is that the DS is present on CP5;

6. The method of carbonation process (CP5) proposed and executed to control the water content during carbonation curing that consists of PDS, WI followed by CC cycles and DS, is a promising way to control the water content on specimens during the carbonation curing process;
7. The highest compressive strength was achieved on the specimen TMWM00C-CP2, which reached 29.7 MPa, followed by TMWM50C-CP2, which reached 26 MPa.

Therefore, based on the compressive results obtained in the current work, the magnesium oxide-rich powder (MRP) used is a promising cementitious binder material in which the compressive strength is related to the carbonation curing process. Besides that, it may represent an alternative way to make hybrid binders containing other mineral powders materials, such as tungsten mining waste mud (TMWM). Finally, the binders produced may represent an effective way to incorporate a wide variety of wastes, and at the same time, capture and store CO$_2$ into cementitious materials. However, future studies are necessary to understand better the binders produced in this work.

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