Hydraulic binder obtained from recycled cement and sand powder

Abstract

Environmentally eco-efficient concrete is currently a popular choice of construction material. Reviewing relevant literature indicates that heating of cement paste can remove hydrating water, and that quartz sand of appropriate granulometry can act as a supplementary cementitious material; however, there have been virtually no reports on the reuse of these materials as alternative binders. This work evaluates the phases present in cementitious compounds produced with binders obtained by thermomechanically treating cement pastes with and without a 15% substitution of quartz sand powder (specific surface area 37.4 m²/g), referred to as RCQ and recycled cement RC, respectively. Testing included thermogravimetric analysis (performed with stripped gases), X-ray diffraction, and scanning electron microscopy with phase identification. The obtained data indicated RC cement paste possesses a predominantly ettringite microstructure, whereas the presence of calcium hydroxide and hydrated calcium silicate predominates in RCQ paste.

Keywords: construction residues, sand functionalization, cement microstructure, thermal analysis, X-ray diffraction.

Resumo

Atualmente tem-se buscado a obtenção de concretos ecoeficientes por meio da reciclagem dos resíduos de concreto. Estudos da literatura mostram que o aquecimento da pasta pode remover a água de hidratação e que areia quartzosa pode atuar como material cimentício suplementar quando na granulometria adequada. Entretanto, praticamente inexistem relatos sobre reuso desses materiais como aglomerantes alternativos. Este trabalho avalia as fases presentes em compostos cimentícios confeccionados com aglomerante obtido do tratamento termomecânico de pastas com 15% de pó de areia quartzosa (RCQ) e sem substituição (RC) com superfície específica de 37.4 m²/g. Foram realizadas termogravimetrias com análise de gases desprendidos, difração de raios X e microscopia eletrônica de varredura com identificação de fases. Os dados obtidos mostram que a pasta RC possui microestrutura predominante de ettringita enquanto na pasta RCQ predomina a presença de hidróxido de cálcio e silicato de cálcio hidratado.

Palavras-chave: análise térmica, difração de raios-X, funcionalização de areia, microestrutura, resíduos de construção.
1. Introduction

Hardened Portland cement paste is composed primarily of hydrated calcium silicate, calcium hydroxide, calcium sulfoaluminates, non-hydrated cement particles, and water (Taylor et al. [1]). According to Mehta and Monteiro [2], cement paste contains chemically combined, capillary, adsorbed, and interlamellar water, the latter three types which can be lost by drying and may cause material shrinkage. Chemically combined water, however, is only released when hydrated products decompose because of heating. Taylor et al. [1] analyzed the hydration of Portland cement using X-ray diffraction, thermal analysis, and infrared spectroscopy methods, which are considered satisfactory for describing this phenomenon. Other studies on Portland cement compound hydration levels have been based only on thermal analyses of hydrated-phase decomposition products (Taylor et al. [1], Shui et al. [3]; Lemonis et al. [4]). Various studies have identified the existence of seven different reactions involved in the decomposition of the cementitious paste: five dehydration reactions, one dehydroxylation reaction, and one decarbonation reaction (Fordham and Smalley [5]). The five dehydration reactions are related to chemically combined, interlamellar, capillary, and adsorbed water losses. Kovler and Roussel [6] and Schöler et al. [7] conducted studies on the hydration and dehydration kinetics of Portland cement through different curing processes and additions. The influence of superplasticizers on the hydration process has also been investigated using atomic modeling techniques (Karen et al. [8]). Various studies have addressed the rehydration of cement after high temperature exposures. Farage et al. [9] analyzed the behavior of cement pastes with a 0.5 water-cement ratio (w/c) subjected to temperatures of up to 300 °C and found it was possible to rehydrate both calcium silicates and non-hydrated grains. Other studies have confirmed the recovery of cement phases subjected to temperatures up to 1,400 °C and verified phases are recoverable when subjected to temperatures similar to those used in the clinker process, although their reuse is restricted to mortar applications (Splittergerber and Mueller [10]). Various researchers have evaluated the microstructure of cement pastes after heat exposure (Burgh and Foster [11]), slow heating, and cooling cycles. They cement pastes with 0.4 w/c have been found to lose water-related hydration mass at 750 °C (Alonso and Fernandez [12]). According to Wang et al. [13], cured cement pastes exposed to temperatures of 400 °C maintain a constant pore structure and phase composition, whereas the same cement pastes subjected to 800 °C temperatures may exhibit new hydration products and refined pore structures. Zanni et al. [14] investigated the behavior of crushed quartz in reactive powder concrete, verifying its low reactivity at approximately ambient temperatures. Klimesch and Ray [15] evaluated the effects of ground quartz surface area on the chemical and physical properties of autoclaved cement pastes using larger particles (0.68 m²/g), and verified these particles acted as nucleating agents, accelerating the formation of calcium silicates and calcium hydroxide. Benezet and Benhassaine [16] confirmed samples of quartz powders may exhibit reactivity with calcium hydroxide, provided they have an average size ≤5 μm. Deschner et al. [17] investigated the hydration of Portland cement using quartz powder and siliceous fly ash with similar granulometries. Under these conditions, they verified that quartz did not exhibit pozzolanic behavior when exposed to ambient temperature (23 °C).

Even when considering the reactivity of sand and possibility of cement rehydration, the reuse of cement and quartz sand as alternative binders has not been reported in literature. This study compared the phases present in cement pastes produced from anhydrous Portland cement and binders, obtained from the thermomechanical treatment of Portland cement pastes with and without the presence of ground quartz sand (sand powder). The objective was to analyze the technical viability of using the sand and cement present in mortar residues utilizing thermomechanical treatment.

1.1 Rationale

The overall consumption of Portland cement has increased over the past three decades, a phenomenon primarily attributed to increasing urbanization and industrialization in countries with emerging economies, as well as the versatility and relatively low cost of the binding materials (SNIC - Sindicato Nacional da indústria do cimento [18]). The production of Portland cement, however, negatively impacts environmental concerns such as natural resources and the climate, and contributes substantially to CO₂ emissions as these are an intrinsic part of its production process. Partial clinker/cement substitutions, particularly those utilizing construction, industrial, and agricultural residues, could thus function as an environmentally efficient concrete alternative.

This study investigated the possibility of reusing dehydrated cement paste and quartzous fine aggregate products, both of which are present in concrete residues, to obtain cementitious and supplementary cementitious materials. Such an approach may contribute to the environmentally friendly disposal of these residues, and could also reduce the environmental impact and carbon dioxide emissions of cement production.

2. Materials and methods

This analysis utilized Portland cement, with a high initial strength

Table 1

| Initial setting times (min) | Final setting times (min) | Loss on ignition 1,000 °C (%) | Loss on ignition 500 °C (%) | Insoluble residue (%) | Specific surface area (Blaine) (m²/kg) | Specific mass (g/cm³) | Compressive strength (MPa) |
|---------------------------|--------------------------|-----------------------------|----------------------------|-----------------------|--------------------------------------|----------------------|--------------------------|
| 131                       | 185                      | 3.5                         | 0.7                        | 0.8                   | 470.8                                | 3.11                 | 30                       | 46                       | 55                       |

1179
and low additive content; anhydrous cement (R), shown in Tables 1 and 2; potable water; and quartz sand powder, produced by grinding to achieve an average particle size of 3.6 µm (Table 3). Grinding was performed in a high energy planetary mill (Planetary Mill Pulverisette 5) at 300 rpm using grinding vessels and zirconium oxide balls. Cement fineness was evaluated by measuring the specific surface area using a Blaine permeability apparatus. The specific surface areas of sand samples were determined using the linearized Brunauer, Emmett and Teller (BET) method in the relative pressure range of 0.05–0.30, achieved with branded equipment from Quantachrome Instruments (Nova Station A-1200e Surface Area and Pore Size Analyzer). Nitrogen gas was used as an adsorbate during testing. Samples were treated at 200 °C for 5 h prior to contact with the gas.

Experiments were conducted in two stages. In the first one, temperatures most suitable for recovering the anhydrous phases of cements were determined by means of thermal analysis in the DTG-60H Shimadzu apparatus (atmospheric air and heating rate of 10 °C/min). Portland cement paste of 0.5 w/c, comminuted after a 28-day water immersion curing, was used for this purpose. Considering the obtained results, cement paste samples were subjected to temperatures of 200, 500, 550, and 700 °C in the same equipment used for thermogravimetric analysis (TGA). These samples were analyzed by X-ray diffraction (XRD) with the Philips-PANalytical PW 1710 equipment, using CuKα radiation, 40 KV power, and an electric current of 30 mA.

In the second experimental stage, Portland cement pastes (AC pastes) were produced with the same parameters as those in step 1. Some samples were ground to achieve a specific surface area of approximately 500 m²/kg, similar to that of the experimental cement. This grinding process was conducted using the same equipment as was used to comminute the sand. The material was immediately heated at 550 °C (see first stage data) for 1 h inside a Flyevier muffle furnace (model FE50RPM) equipped with a temperature controller to eliminate water content (free and combined) from the pastes and recover anhydrous cement phases. The duration was determined as a function of the furnace thermal power, cement paste mass, and specific heat to implement an efficient material dehydration process. The binder (O), obtained following thermomechanical treatment, was used to produce new cement pastes by replacing 15% of the thermally treated material with sand powder (RC pastes) and also without replacement (RCQ pastes). Table 4 presents the percentages of water (W), anhydrous Portland cement (R), recovered binder (O), and ground sand used in each sample.

The phases present in the AC, RC, and RCQ pastes were identified using simultaneous thermogravimetric (TG) and mass spectrometry (TGMS) analyses from a PERKIN-ELMER STA-6000 simultaneous thermal analyzer coupled with a HIDEN HPR 20 quadrupole mass spectrometer. These tests were conducted at a heating rate of 10 °C/min under helium flow (20 mL/min). X-ray diffraction was also tested using the same equipment and under the same conditions as those applied in stage 1. The cement paste microstructures were analyzed via scanning electron microscopy with phase identification (SEM-FEG) in a Quanta FEG 3D FEI device with 0.8 nm resolution, 3–99 mm focal length, and a 24–800,000-fold increase in high vacuum.

3. Results and discussions

3.1 Evaluation of recovery temperature of anhydrous phases

3.1.1 Thermal analysis

Figure 1 shows the thermal analysis of the AC paste following comminution, presenting the differential thermal analysis (DTA), TGA, and TGA-derivative curves (DrTGA). These curves are similar to the thermal analysis curves described in literature investigating the hydration of Portland cement pastes (Kovler and Roussel [6]; Ramachandravan [19]; Kakali et al. [20]; Alarcon Ruiz et al. [21]; Chen et al. [22]). Three regions of significant heat and mass variations were observed. The first mass loss, occurring between 30–200 °C, resulted from the dehydration reactions of several hydrates (carboaluminates, ettringite, etc.) including, primarily, calcium silicate hydrate (C-S-H) (Taylor et al. [1]). The second largest mass loss was observed between 400–550 °C, corresponding to the dehydroxylation of Portlandite (Alarcon Ruiz et al. [21]). The third mass loss appeared at approximately 650 °C, where Alonso and Fernandez [12] approximated the decomposition of calcite (CaCO₃) and generation of CO₂ occurred. Furthermore, calcium carbonate experiences decarbonization above 700 °C (Alonso and Fernandez [12]; Rodrigues et al. [23]; Fuentes et al. [24]). Relevant literary information indicates that it is possible to lose the water mass responsible for cement hydration at temperatures of approximately 550 °C.

3.1.2 X-Ray diffraction

The cement paste diffractograms taken at ambient temperature (26 °C) were subsequently heated at 200, 500, 550, 700, and 750 °C, as presented in Figure 2. Calcium hydroxide (P), calcium

| Table 2 | Chemical characterization of Portland cement |
|---------|---------------------------------------------|
| SiO₂ (%) | Al₂O₃ (%) | CaO (%) | SO₃ (%) | CO₂ (%) | C₃A (%) |
| C20 | C20 | C20 | C20 | C20 | C20 |

| Table 3 | Chemical characterization of Portland cement |
|---------|---------------------------------------------|
| Specific surface area (BET) (m²/kg) | Mean diameter (µm) | Pozzolanicity – Chapelle modified (mg/g CaO) |
| 37,400 | 3.6 | 958.2 |

| Table 4 | Cement paste composition: AC, RC, and RCQ pastes |
|---------|---------------------------------------------|
| Paste | W (%) | R (%) | O (%) | Ground sand (%) |
| AC | 33 | 67 | 0 | 0 |
| RC | 33 | 0 | 67 | 0 |
| RCQ | 33 | 0 | 57 | 10 |
carbonate (C), tricalcium silicate (A), dicalcium silicate (B), calcium aluminum hydrosulfate (E), tetracalcium sulfoaluminate (F), and calcium silicate hydrate (H), constituents commonly found in Portland cement pastes (Alonso and Fernandez [12]; Shui et al. [25]), were present in unheated samples. After heating to 200 °C, calcium hydroxide (P), tricalcium silicate (A), calcium silicate hydrate (H), and calcium carbonate (C), were identified. At 500 °C, the identified phases were calcium hydroxide (P), tricalcium silicate (A), and calcium carbonate (C). Calcium hydroxide (P), tricalcium silicate (A), dicalcium silicate (B), calcium carbonate (C), and tettracalcium sulfoaluminate (F) were observed at 550 °C. Crystals of tricalcium silicate (A), calcium carbonate (C), and calcium oxide (O) were identified in paste heated at 700 °C. Existing literature indicates calcium silicate hydrate, calcium oxide, and dehydrated calcium sulfoaluminate coexist at 700 °C (Schöler et al. [7]; Splittgerber and Mueller [10]; Janotka and Nürnbergerov [26]). It was noted that calcium silicate decomposed at 750 °C into new phases, reducing the mechanical strength of the material (N) (Splittgerber and Mueller [10]; Vieira et al. [27]; Trezza and Lavat [28]).

All cement pastes demonstrated peaks illustrating the presence of calcite, indicative of carbonation. Hydrated calcium silicate was observed in two samples: unheated, and thermally treated at 200 °C. Based on these results, thermal analysis data, and existing literature, it was determined that cement pastes treated at temperatures >700 °C may contain calcium silicate hydrates decomposed into new phases (Splittgerber and Mueller [10]; Vieira et al. [27]; Trezza and Lavat [28]). Because of this, the 700 °C temperature point was taken as a reference for anhydrous cement recovery. However, pastes produced using cement obtained by thermal treatments at this temperature exhibited insufficient cohesion for molding (Splittgerber and Mueller [10]; Trezza and Lavat [28]). Conversely, pastes produced using cement that underwent 550 °C thermal treatment exhibited satisfactory cohesion levels. A working temperature of 550 °C was consequently selected for binder recovery. Moreover, the suitability of this temperature was verified by comparing the diffractograms of cement pastes treated at 550 °C (recovered binder) to those of Portland cement (Figure 3). Both diffractograms contained the same main peak phases: calcium hydroxide (P), tricalcium silicate (A), dicalcium silicate (B), calcium carbonate (C), and calcium ferroaluminate (F).

3.2 Investigation of cement pastes produced using recovered cement with and without sand powder

3.2.1 Thermogravimetric analysis

Below, the thermogravimetric analysis results of three pastes are

![Figure 1](https://via.placeholder.com/150)

**Figure 1**

Thermal analysis of comminuted cement paste (w/c 0.5)

![Figure 2](https://via.placeholder.com/150)

**Figure 2**

Diffractograms of cement pastes at ambient temperature (26 °C) and thermal treatments: 200, 500, 550, 700 °C, and 750 °C. Labeled peaks: tricalcium silicate (A), bicalcium silicate (B), calcium carbonate (C), calcium aluminum hydrosulfate (E), tetracalcium sulfoaluminate (F), calcium silicate hydrate (H), calcium hydroxide (P), new silicate (N), calcium oxide (O)

![Figure 3](https://via.placeholder.com/150)

**Figure 3**

Diffractograms of Portland cement and recovered binder (paste thermally treated at 550 °C). Peaks: calcium hydroxide (P), tricalcium silicate (A), dicalcium silicate (B), calcium carbonate (C), calcium ferroaluminate (F)
presented: a paste produced from Portland cement, one from re-
covered cement without partial replacement, and one with a par-
tial replacement (15% quartz sand powder). The obtained curves,
shown in Figure 4, are similar to those described in relevant litera-
ture where thermal analysis was used to investigate the hydration
of Portland cement and Portland cement paste without high addi-
tion levels (Fordham and Smalley [5]; Farage et al. [9]; Alonso and
Fernandez [12]; Vieira et al. [27]; Zhutovsky and Kovle [29]; Hoppe
Filho et al. [30]). Observations indicate high mass losses at three
critical temperatures, which synthesized the dehydration process
and resembled the results obtained in step 3.1.1. The first mass
loss occurred between 100–200 °C as a result of the dehydration
reactions of various hydrates (Farage et al. [9]). The second sig-
nificant mass decrease was observed between 400–550 °C, corre-
sponding to the dehydration of Portlandite (Fordham and Smalley
[5]; Rodrigues et al. [23]; Zhutovsky and Kovle [29]; Hoppe Filho
et al. [30]). The third mass loss occurred at approximately 750 °C,
corresponding to the decarbonization of calcium carbonate pro-
duced by the decomposition process of calcium silicate hydrate (C-
S-H) and leading to the production of calcium oxide (CaO), which
in turn reacted with CO₂ in the atmosphere of the equipment to
produce calcite (Alonso and Fernandez [12]; Chen et al. [22]; Shui
et al. [25]; Zhutovsky and Kovle [29]).

Comparison indicated the RC paste experienced a greater mass
loss than the RCQ paste, potentially associated with the ettr-
ingite formation resulting from interactions between calcium,
sulfate, aluminate, and hydroxyl ions present in the phases
identified in the cement after exposure to 550 °C temperatures
(Mehta and Monteiro [2]). The lower mass loss in RCQ paste
was identified in the Portlandite decomposition region, indicat-
ing a lower amount of calcium hydroxide was present and dem-
onstrating the pozzolanic behavior of the sand powder (Benezet
and Benhassaine [16]).

### 3.2.2 Mass spectrometry

Figure 5 shows the data obtained during the TGA of the volatile
products released by the AC, RC, and RCQ pastes, respectively.
This data indicates two types of gases were produced when the
pastes were subjected to heat: one containing hydrogen and oxy-
gen at a temperature of approximately 450 °C (blue curve), and
another containing carbon and hydrogen in the temperature range
of 580–750 °C (green curve). Considering the present phases, it
was deduced one of these gases was water in the gaseous state,
resulting from dehydration, whereas the other was CO₂ gener-
ated by the decomposition of calcium carbonate (Fordham and
Smalley [5]; Ramachandran [19]; Rodrigues et al. [23]). These
gases were produced by all samples at temperatures of approxi-
mately 450 and 720 °C. Peak observations verified the amount
of the produced CO₂ was the same in the three samples, AC,
RC and RCQ. However, this amount was significantly reduced
in the samples of thermally treated cement and cement partially
replaced by sand powder. This indicates a lower amount of de-
hydroxylation occurred in the Portlandite reaction, and that sand
powder possesses lower pozzolanicity.

### 3.2.3 X-ray diffraction

The AC, RC, and RCQ paste diffractograms are shown in Figure 6.
The phases present in the AC paste are the same as those identi-
fied in the RC paste (made from thermomechanically treated ce-
mlet). The primary identified crystalline constituents were calcium
carbonate (C), calcium hydroxide (P), hydrated calcium silicate
(H), dicalcium silicate (B), and tricalcium silicate (A). An additional
peak related to the presence of silicon oxide (Q) in crystalline form
was identified in the RCQ paste (produced using a binder obtained
via thermomechanical treatment and partial sand powder replace-
ment) (Bacarji et al. [31]).

Figure 5
Volatile products released by pastes produced with anhydrous Portland cement (AC); recycled cement
(RC), and 15% replacement by sand powder (RCQ)
3.2.4 Microstructural analysis

Figure 7 shows the typical AC, RC and RCQ paste microstructures. During microstructural analysis with phase identification (MEV-FEG), calcium hydroxide and hydrated calcium silicate were observed in all pastes. The presence of ettringite predominated in the RC paste image, whereas ettringite was not abundant in the AC paste image, nor present in the RCQ paste. This phenomenon may be associated with the reduced formation of ettringite caused by the interactions between calcium, sulfate, aluminate, and hydroxyl ions with water (Mehta and Monteiro [2]), and could be explained by the reaction between the sand powder and constituents of the recovered cement when subjected to hydration. This hypothesis reinforces the theory of the supplementary behavior of sand power in cement, as well as the reactivity increase caused by its comminution (Rêgo et al. [32]).

4. Conclusions

- Pastes made with thermomechanically treated cement (550 °C and comminution) without replacement by sand powder presented primarily ettringite microstructures.
- Pastes made with heat-treated recycled cement and cement partially replaced by sand powder exhibit significantly lower amounts of gas release as compared to pastes made with CPV-ARI Portland cement or heat-treated recycled cement without sand powder substitutions, indicating the dehydroxylation reaction occurs to a lesser extent in Portlandite and the pozzolanicity of sand powder was relatively lower.
- Pastes made with cement recovered by thermomechanical treatment (550 °C and comminution) and partial (15%) sand powder (specific surface area of 37 m²/g) replacements, demonstrated similar phases to those found in CPV-ARI Portland cement, indicating it is possible to recover cements that have been subjected to this temperature.
- The simultaneous use of cement from thermally treated pastes and sand powder from quartz sand grinding demonstrated the technical viability of recovering cements and sands present in mortar and concrete residues via thermomechanical treatment.
- This study represents the first step in evaluating the reuse of cement and sand obtained from construction wastes as alternative binders; however, further complimentary research is needed.

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