Bentonites in Southern Spain. Characterization and Applications

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Abstract: The objective of this work was to investigate and demonstrate the pozzolanic properties of the bentonites found at the San José–Los Escullos deposit, located in the southeast of the Iberian Peninsula, to be used in the manufacturing of more durable and environmentally compatible pozzolanic cements, mortars and concretes. These bentonites are mainly composed of smectites, with montmorillonite as the main clay mineral. They were formed by the hydrothermal alteration of tuffs, volcanic glasses, dacites, rhyolites and andesites. For this research, samples were taken from outcrops on the south, north and west side of the San José–Los Escullos deposit, and in the Los Trancos deposit located 19.3 km to the northeast. All samples consisted of bentonites, except for a zeolite sample taken from the northern flank of the San José–Los Escullos deposit, which was used to contrast and compare the behaviour of bentonite in some of the analyses that were done. An investigation of the mineralogical, petrological, chemical and thermogravimetric characteristics of the samples was carried out using various methods, such as XRD, OA (Oriented aggregates), TGA, XRF, SEM and thin section petrography (TSP). In addition, a chemical analysis of pozzolanicity (CAP) was done at 8 and 15 days to determine the pozzolanic capacity of the samples. XRD, XRF, SEM and TSP studies showed that these bentonites have a complex mineralogical constitution, composed mainly of smectites of the montmorillonite variety, as well as halloysite, illite, vermiculite, biotite, muscovite, kaolinite, chlorite, mordenite, feldspar, pyroxene, amphibole, calcite, volcanic glass and quartz. Thermogravimetric analysis (TGA) established the thermal stability of the bentonites studied at above 800 °C. Chemical analysis of pozzolanicity (CAP) confirmed the pozzolanic character of the bentonites, exhibited in their reactive behaviour with Ca(OH)₂. The pozzolanic reactivity increased significantly from 8 to 15 days. These results show that the materials studied can be used as quality pozzolans for the manufacture of pozzolanic cements, mortars and concretes.

Keywords: bentonite; mordenite; smectite; pozzolanicity test; cement; concrete

1. Introduction

This research focused on the characterisation and uses of bentonites from the San José–Los Escullos deposit as pozzolans for the manufacture of pozzolanic cements, mortars and concretes. There are no previous reports on the use of bentonites from this deposit for these specific purposes, so the results presented here are considered new. However, the use of bentonites is broad and diverse, as described below.

Bentonites have been used in many fields for several decades. Due to their chemical, mineral and technical properties, the bentonites from Sardinia (Italy) were used to prepare bentonite pastes to be used in pelotherapy [1], thus substituting traditional peloids. Nowadays, the use of bentonites is so diverse that they are used in many sectors of the economy, science and technology [2]. There is a current trend to thermically activate the bentonites to facilitate the production of geo-polymer binding agents [3]. The heated bentonites quickly acquire pozzolanic activity, as has been proven by Habert et al. [4]. They proved that by heating the bentonites between 20 and 900 °C it is possible to raise the mechanical resistance...
of the mortar up to 60 MPa in 28 days. Kaci et al. [5] proved the influence of bentonite on
the rheologic behaviour of the mortar, especially regarding the fluency and thixotropy, and
arrived at the conclusion that bentonite improves the resistance of the mortar. To mitigate
the effects of copper mining, Bertagnolli et al. [6] used thermically activated bentonitic
clay in the porous beds to absorb the copper. Slamova et al. [7] proved the role of clay
in animal food and formulated procedures of application to avoid gastrointestinal disorders
in these animals; they highlighted that the clay minerals with a fine granulometry and a
high capacity for absorption are the most effective for veterinarian use. Pelayo [8] proved
the behaviour of bentonite as an efficient insulator in storing radioactive residue in his
research on bentonites from the Morrón de Mateo mineral deposit, in the south east of the
Iberian Peninsula. Hassan and Abdel [9] made a mixture out of bentonite–zeolite with a
1:10 ratio to improve the productivity of the crops in sandy and arid soils. Recent research
shows great appreciation for products that are absorbent, catalysts and biomaterials [10].
Park et al. [11] list a series of uses for the clay bentonites in drilling mud, whitening prod-
ucts, stabilising emulsions, in the smelting industry, drying products, catalysts, adhesive
seals, cosmetics and pharmaceutical products. Panday and Ramontja [12] list in detail the
most recent progress in the use of bentonite and its compounds to eliminate contaminating
synthetic dyes in water. Luqman and Enobong [13] used bentonite as a base to prepare
sol-gel to obtain bioactive glass, significantly lowering the costs generated using old pre-
cursors to alkoxy silanes. Aravindraj and Sapna [14] were able to raise the resistance to
compression and flection of concrete, mortars and cements, substituting the cement for
bentonite in 15%, which also made it exceptionally durable against the effects of sulphate
and chlorides. In the process of de-fluoridisation of the waters, Masindi [15] patented a
simple, new and innovative method in the synthesis of cryptocrystalline magnesite from
bentonites, which notably surpasses the absorption capacity of fluoride in conventional
products. Masindi [15] also proved that bentonite that is ground up to the correct size
can be used efficiently in the management, neutralisation and remediation of acid waters
from mines. The effectiveness of this fact is because the thermic activation of the pores
and micropores increases, thus facilitating an ionic exchange between the copper and the
interlamellar cations. Kim et al. [16] have recently proposed a numerical model based on the
relationship between density and thermal–hydraulic properties of compacted bentonite for
sealing high-activity waste tanks. Masood et al. [17] made compound concretes with low
calcium and arid bentonite compounds that came from recycled concrete. The bentonite
substituted the cement in 5%, 10%, 15% and 20%. The concretes were significantly efficient
regarding durability and mechanical resistances; on the other hand, it was highlighted as
a good response to the environmental problem. Wu et al. [18] made a mixed compound
based on reactive MgO, blast furnace grinded granulated slag, bentonite and clay sand
to improve soils. Bentonites are currently being used successfully to make high activity
radioactive waste inert, in the form of a paste fabricated from grains, particulates and
bentonite dust [19].

2. Geological Settings

The study area is in the province of Almería (Spain) between the municipalities of San
José de Níjar and Los Escullos (Figure 1). The research took place inside of the Los Frailes
caldera, a volcanic structure that was formed 14.4 million years ago in the southeast of the
Iberian Peninsula, before the other surrounding calderas were formed, as Rodalquilar and
La Lomilla. The Los Frailes caldera is a circular megastructure slightly larger than 5 km in
diameter [20,21].

The geology is made up of rocks that were formed during the volcanic processes of the
Neogene and are mainly andesites and dacites from the Unidad Frailes 1 (UFR-1), dacites
and rhyolites from the Rodalquilar Complex, as well as basaltic andesites and post-caldera
pyroxene andesites from the Unidad Frailes 2 (UFR-2) [21–23]. The rocks of a half acid
composition, as dacites and rhyolites, make dykes and domes, whereas the most basic
lithologies like the andesites, form great lava flows and pyroclastic breccia (Figure 2) [24].
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Generally, bentonites are stratified, banded and changed by spheroidal weathering. They are highly friable and mylonitised. Their colour is a characteristic beige (Figure 3). Occasionally they include dacite pyroclastic and tuff materials. The outcrops are cut by vertical and subvertical diaclases with a dip between 70° and 80° towards the west [24].
The bentonite deposits are well represented in the study area. In UFR-1, the bentonite has been formed by fine lapilli and bentonised ash; these are in Cerro de la Palma and in Barranco de Cala Figuera. In Rodalquilar, the dacite tuffs are bentonised and form industrial deposits in the Rambla del Plomo [8,25–27].

The most representative outcrops of the area of research are in a small area of the southern flank of the San José–Los Escullos deposit, as well as in the western part of the same area (Figure 3a,b).

3. Materials and Methods

3.1. Materials

The research area is in the east of San José and to the west of Los Escullos. In this place the bentonites form a representative outcrop on the southern side of the San José–Los Escullos deposit. Five samples of bentonite were taken directly from the outcrops for this research and were designated with the codes B-01 through B-05 (Figure 4a).

Another three samples were taken outside of the deposit limits to make comparative studies. These are: BFO-01, ZFN-01 and BLT-01 (Figure 4a,b).

The dacite has a massive, uniform, vesicular and brecciated structure. It has a medium-grained, porphyritic, phaneritic texture. Its colour varies from dark grey to green. It is generally altered by limonite, sericite and chlorite. The dacite is composed of phenocrystals of amphibole, pyroxene and plagioclase, as well as sericite, glass, iron oxide and opaques.
Andesite is dark grey to green in colour, with a brecciated and prismatic-columnar structure. It is composed of plagioclase, pyroxene, glass, sericite and opaques. Its texture is porphyritic. It is strongly altered to zeolite near the fault systems. Plagioclases are sericitised, while pyroxenes are chloritised. The pyroxenes and plagioclases are replaced by zeolites.

Bentonite varies in colour from light grey, green, beige to yellow. It has a stratified structure. The texture is cinereous and devitrified. It is formed mainly by smectite, and montmorillonite the main mineral, as well as altered glass, relict crystals and fragments of rocks altered to smectite, zeolite and serpentine. The bentonite is weathered and fractured in the outcrops.

Zeolite is white, light grey and green in colour. Its structure is massive and compact, becoming brecciated. It is mainly composed of the mordenite, accompanied by smectite, plagioclase, potassium feldspar, quartz, hematite, amorphous materials and traces of gypsum. They are altered in outcrops, forming shallow, residual and porous clay layers. Outcrops are marked by abundant diaclases and fissures. Mordenite appears as elongated, fibrous and tabular crystals associated with montmorillonite, halloysite, kaolinite and micas.

3.2. Methods

3.2.1. Petrography Characterisation by Thin Section (PTS)

Petrographic analysis was carried out to describe the properties of the samples analysed: mineral type, texture, colour, grain size, alterations (zeolitisation and bentonisation) as well as petrogenesis. The equipment used was a Leica DM600M microscope (Leica, Wetzlar, Germany). It has an adapted system of 13 filters (DTA-RPFMAX) for visible and infrared light at 13 wavelengths, from 350 nm to 1000 nm, in 50 nm intervals. It has a LAS control and a Märzhäuser motorised plate. The image analysis software used was APHELION.

3.2.2. X-ray Diffraction (XRD)

In the sample analyses the Phillips diffractometer 1710 BASED (Amsterdam, Netherlands) was used with the following characteristics: tube anode of Cu, generator tension (45 Kv), generator current 30 mA, wavelength Alpha 1 [Å] 1.54060, wavelength Alpha 2 [Å] 1.54439, intensity ratio (alpha1/alpha2) 0.500, scan range of software 2 theta (4–60°), monochromator, and PC-APD diffraction software.

3.2.3. Oriented Aggregates (OA)

This method was used to determine the clay minerals associated with more crystalline species, as feldspars, quartz and zeolites, present in the researched samples. Some types of clay, such as smectites, chlorites and vermiculites have a d = 14 Å. The test procedure consisted of taking 500 mg of each bentonite sample previously crushed, ground and sieved to 2 µm. Next, the samples were placed in test tubes and mixed with 4–5 drops of sodium hexametaphosphate. They were then allowed to settle. A part of the saturated clay suspension was placed in a glass sample holder for 24 h at room temperature. After evaporation of the water, a thin layer of clay remained on the sample holder, which was treated with ethylene glycol and heated in an oven at 70–80 °C for 24 h. During this time, the vapours acted directly on the sample. The samples were then analysed by XRD and the corresponding patterns were obtained.

3.2.4. Thermogravimetric Analysis (TGA)

This method consisted in heating the samples and a pattern in a controlled manner. During this process, the temperature difference (ΔT) was measured based on time. The thermogravimetric analysis (TGA) measured the variation and behaviour of the mass of the samples under the influence of temperature in a controlled atmosphere. This variation manifested in mass loss. The recording and monitoring of the changes gave information
regarding the decomposition of the sample or of its reaction to other components. This test used Universal equipment V4.1DTA Instruments/2960 SDT V3.OF. A total of 13.1971 g was analysed for each sample using the 10C-1100N2, N2 10 mL/min method, and the TGA curves were obtained.

It was important to apply this method to the samples because it was important to know the sorbed phases (adsorbed, absorbed and constituent water), porosity, reactions that occur with increasing temperature, weight loss, variations in cation exchange capacity (CEC), increase in active surface area, thermal stability and pozzolanic capacity. These properties are decisive in the manufacture of pozzolanic cements mixed with natural and activated bentonites, which is highly important for the preservation of the environment and the durability of sustainable building materials.

3.2.5. X-ray Fluorescence (XRF)

This research used a Philips PW 1404 (Amsterdam, Netherlands) with a collimator, with a radiation intensity between 10 and 100 kV, and monochrome was used to isolate the radiation. The preparation of the samples consisted of a previous grinding to 200 mesh. Next, 8 g of dust with a 1.5 mL of elbaite were mixed in to make a 5 cm in diameter paste to be analysed by XRF.

3.2.6. Scanning Electron Microscopy (SEM)

The following equipment was used to prepare and analyse the samples: Hitachi S-570 Scanning Electron Microscope (Tokyo, Japan) with a KeveX 1728 analyser, the BIORAD Polaron Division Carbon Evaporation Power Supply and the Polaron SEM Coating System. The Winshell and Printerface software were also used. All the samples were reduced to between 0.2 and 0.5 cm; and were then covered by a layer of graphite and placed in the sample carrier for their analysis.

3.2.7. Chemical Pozzolanicity Analysis (CPA)

Pozzolanicity is the property of rocks and minerals to actively react with calcium hydroxide at 40 degrees, once crushed, ground up and micronised to around 63 µm. Under these conditions they can acquire cementitious properties and develop mechanical strength over time [29]. The aim of this work is to determine the pozzolanic behaviour of the analysed bentonites and to establish their uses in the manufacture of cements, mortars and concretes with excellent durability and mechanical strength.

Pozzolanicity was determined by comparing the amount of calcium hydroxide that exists in a watery dissolution which contains hydrated cement and pozzolan, with enough calcium hydroxide necessary to obtain a saturated dissolution with the same alkalinity as the other sample. The chemical trial for pozzolanicity was done following standard UNE-EN 196-5:2011 [29] to 8 to 15 days. An amount of 100 mL of distilled water was heated to 40 °C, for 60 s. Twenty grams of sample (bentonite) and cement made up of 75:25 ratio was added. After a week, the solution was filtered. The concentration of the hydroxyl ions [OH\textsuperscript{−}] was calculated with the following equation:

\[
[\text{OH}^-] = \frac{1000 \times 0.1 \times V_3 \times f_2}{50} = 2 \times V_3 \times f_2
\]

where:
- [OH\textsuperscript{−}]: is the concentration in hydroxyl ions (mmol/L).
- \(V_3\): is the volume of the hydrochloric acid solution (0.1 mol/L).
- \(f_2\): is the factor of the hydrochloric acid solution (0.1 mol/L).

The concentration of calcium oxide (CaO) was calculated using the following equation:

\[
[\text{CaO}] = \frac{1000 \times 0.025 \times V_4 \times f_1}{50} = 2 \times V_4 \times f_1
\]
where:
• \([\text{CaO}]\): is the concentration in calcium oxide (mmol/L).
• \(V_4\): is the volume of EDTA solution used in the titration.
• \(f_1\): is the factor of the EDTA solution.

The pozzolanicity test is positive when the hydroxide concentration of calcium in dissolution is lower than the saturation concentration [29]. The results are seen in the graph, where all samples below the solubility isotherm curve are considered as pozzolans.

4. Results and Discussion

4.1. Petrographic Study by Thin Section

The analysed bentonites have a texture that is porphyric, relict, hemivitreous, crystal-lithoclastic, pyroclastic and oriented. The predominant minerals are clays (montmorillonite-type smectite), mordenite, plagioclase, biotite, muscovite, hematite, apart from the glass. Subordinately there is quartz, kaolinite and polygenic lithic fragments of a small diameter. Figure 5a,b (N//) represents a process of strong pyroxene and plagioclase alterations being substituted partially or totally by smectite and mordenite.

The micrograph of the sample BLT-01 (Figure 5c) (Nx) shows a predominance of smectite and muscovite amid transforming into smectite. The hematite is in small fissures and covers the smectite and the mica. Figure 5d (Nx) shows fractured pyroxene phenocrystal, completely altered to zeolite, floating on a glass fabric. The biotite crystals show a reaction with the fabric and are partially surrounded by halos of hematite in the paste. There is a biotite crystal in process of matrix alteration (Figure 5a,c,d). Three types of matrices can be distinguished: in sample B-01, zeolitised; in B-05, combined (zeolitised–bentonised); in BLT-01, bentonised; and in ZFN-01, glassy, with zeolite formation. The comparison among the samples analysed shows that the bentonisation process was more effective in sample BLT-01.
BLT-01 (Figure 5c), with total development of the clay minerals. In the case of samples B-01, B-05 and ZFN-01, bentonisation did not totally alter the pyroxenes, amphiboles, plagioclase, biotite and muscovite, as these minerals still persist. This fact is observed in Figure 5a–d.

4.2. X-ray Diffraction

The phase study showed that the samples analysed have a complex and heterogeneous mineralogy, with the exception of sample BLT-01. The following minerals were detected: montmorillonite-type smectite, illite, vermiculite, halloysite, biotite, muscovite, mordenite, orthoclase, plagioclase, hematite, kaolinite, chlorite, calcite and quartz (Figure 6). According to Figure 6, each sample behaves according to its mineral content. Comparison of X-ray diffraction patterns reveals a clear similarity between samples BLT-01, BFO-01 and B-05, suggesting the presence of similar mineralogical phases. Something very different is observed in samples B-01 to B-03, which shows a greater complexity than the other samples. This is due to the presence of mordenite, plagioclase, muscovite, quartz and kaolinite in greater quantities. The presence of smectite in sample BLT-01 is more evident than in samples BFO-01 and B-05, while in B-01 to B-03, there is a greater abundance of mordenite.

Figure 6. X-ray diffraction patterns of bentonite from the research area.
According to the abovementioned and to the position of the samples in the deposit (Figure 4), it can be deduced that the bentonisation process was less intense towards the centre of the San José–Los Escullos deposit. Samples B-01 to B-03 represent another case of typical zonality in this deposit, as they indicate that the zeolitisation process, unlike bentonisation, decreases from the centre towards the flanks.

The presence of orthoclase, kaolinite and quartz could indicate a previous deposition of acidic volcanic material which was altered to form mordenite and smectites. The presence of hematite suggests that hydrothermal activity took place in this region. This interpretation agrees with Costafreda [24].

The bentonisation and zeolitisation process in the south and southeast of Spain was practically ubiquitous, as has been stated by Martín-Vivaldi [30]; Leone et al. [26]; Linares [31]; Delgado and Reyes [32]; García-Romero et al. [33]; and Martínez et al. [34].

Samples BFO-01 and BLT-01 lie to the west and south of the San José–Los Escullos deposit, respectively. However, the chemical composition (Table 1), as well as the mineralogy, indicate that both bentonisation and zeolitisation processes were caused by large-scale hydrothermal alteration of materials of calc-alkaline composition.

### Table 1. Chemical composition of bentonites from the San José–Los Escullos deposit and surrounding areas (% weight).

| Sample | SiO₂ | Al₂O₃ | CaO | Na₂O | K₂O | MgO | Fe₂O₃ | TiO₂ | LOI * | Na₂O/CaO | MgO/Al₂O₃ | Si/Al |
|--------|------|-------|-----|------|-----|-----|-------|------|-------|-----------|------------|------|
| B-01   | 64.30| 13.71 | 1.03| 3.3  | 2.5 | 1.3 | 0.123 | 11.5 | 0.58  | 0.18      | 4.1        |      |
| B-02   | 51.29| 13.13 | 1.20| 0.53 | 7.61| 2.75| 0.122 | 22.0 | 1.17  | 0.58      | 3.4        |      |
| B-03   | 52.53| 13.69 | 1.11| 0.99 | 0.70| 7.67| 2.43  | 0.133| 20.5  | 0.89      | 0.56       | 3.3 |
| B-04   | 61.3 | 14.76 | 1.24| 1.56 | 1.92| 5.0 | 2.46  | 0.116| 11.5  | 1.26      | 0.34       | 3.6 |
| B-05   | 62.46| 14.57 | 0.94| 2.64 | 2.21| 4.0 | 1.83  | 0.133| 11.5  | 2.8       | 0.27       | 3.7 |
| BFO-01 | 52.13| 17.05 | 1.12| 1.96 | 0.35| 5.91| 1.61  | 0.123| 19.2  | 1.75      | 0.35       | 2.7 |
| BLT-01 | 47.26| 19.02 | 1.22| 0.30 | 0.87| 4.5 | 2.45  | 0.173| 24.4  | 0.25      | 0.24       | 2.1 |
| ZFN-01 | 67.04| 12.55 | 1.54| 2.64 | 1.79| 1.40| 1.17  | 0.085| 11.7  | 1.75      | 0.11       | 4.7 |

* LOI: Loss on ignition.

### 4.3. Oriented Aggregates

The analysis of the oriented aggregates attenuated most of the more crystalline phases present in the samples, like mordenite, plagioclase, quartz, orthoclase and others; thus, the following clay minerals were clearly observed in the X-ray diffraction patterns: smectite, vermiculite, illite, kaolinite and halloysite (Figure 7).

Figure 7 represents a case where there is an appreciable amount of mordenite in the sample (ZFN-01), taken at the north of the deposit. Initially the peaks of mordenite of greater intensity masked the clay minerals, whereas, once the sample was treated with ethylene glycol, the vermiculite peaks became visible, as well as illite, kaolinite and halloysite (Figure 7).

Another detail worth highlighting is that the main peak of smectite in sample (ZFN-01) is initially in position 2θ = 6.5 with interplanar spacing equal to 14 Å; once the sample was treated with ethylene glycol it produced a strong peak reflection of the smectite and a displacement from 2θ = 6.5 to 2θ = 5.4 (Figure 7). In this case, the peak reflection of the smectite has a maximum intensity of 100% and a maximum spacing of 17 Å. This fact proved that the bentonite of the San José–Los Escullos deposit is expansive.
Delgado and Reyes [32]; García-Romero et al. [33]; and Martínez et al. [34]. Samples BFO-01 and BLT-01 lie to the west and south of the San José–Los Escullos deposit, respectively. However, the chemical composition (Table 1), as well as the mineralogy, indicate that both bentonisation and zeolitisation processes were caused by large-scale hydrothermal alteration of materials of calc-alkaline composition.

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|--------|------|-------|------|------|-----|-----|-------|------|------|---------|-----------|-------|
| B-01   | 64.30| 13.71 | 1.03 | 3.3  | 2.3 | 2.5 | 1.3   | 0.123| 11.5 | 0.58    | 0.18      | 4.1   |
| B-02   | 51.29| 13.13 | 1.20 | 1.40 | 0.53| 7.61| 2.75  | 0.122| 22.0 | 1.17    | 0.58      | 3.4   |
| B-03   | 52.53| 13.69 | 1.11 | 0.99 | 0.70| 7.67| 2.43  | 0.133| 20.5 | 0.89    | 0.56      | 3.3   |
| B-04   | 61.3 | 14.76 | 1.24 | 1.56 | 1.92| 5.0 | 2.46  | 0.116| 11.5 | 1.26    | 0.34      | 3.6   |
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4.4. Thermogravimetric Analysis

The thermal behaviour of the samples analysed shows similarities and differences, according to the variables in the curves (Figure 8). The curve of sample ZFN-01 shows a higher mass loss compared to the other samples. Like Costafreda [24], the authors have linked this fact to the complex composition of the zeolite, due to its Si/Al ratio, its capacity for cationic exchange and the proportion of smectite–mordenite. The BLT-01 sample is noted because of its mass loss, which is coherent with the data of LOI represented in Table 1. The BFO-01, B-02 and B-03 show similar behaviour which may be related to its low ratio of Si/Al, its high values of LOI and the appreciable content of MgO. The samples B-01 and B-04 behave similarly with regards to their chemical composition, high contents of SiO₂ and Na₂O and the lowest values of LOI.

In addition, it is noted that the TGA curves show a simple decomposition in all samples analysed (Figure 8). In all cases there are three significant inflection points that signal different thermogravimetric processes. These changes occur at intervals of temperature 39.2–86.1 °C, 86.1–198.1 °C and 198.1–502.4 °C, which indicate phases of mass loss during heating. Between 39.2 and 86.1 °C the mass loss is 2.362%, and it indicates a loss of humidity and superficial dehydration in the smectite–mordenite phase. Between 86.1 and 198.1 °C, the mass loss is of 3.972%, which points to the continuation of the process of dehydration, loss of superficial humidity and gasses removal. Between 198.1 and 502.4 °C, the curve shows a loss of mass equivalent to 2.291% due to the loss of intrareticular water. At 750 °C the curve stabilises, which could be due to the high Si/Al ratio of the mordenite. At this temperature range, the mordenite collapses and the sample structure is reordered.
Figure 7 represents a case where there is an appreciable amount of mordenite in the sample (ZFN-01), taken at the north of the deposit. Initially the peaks of mordenite of greater intensity masked the clay minerals, whereas, once the sample was treated with ethylene glycol, the vermiculite peaks became visible, as well as illite, kaolinite and halloysite. Another detail worth highlighting is that the main peak of smectite in sample (ZFN-01) is initially in position \(2\theta = 6.5\) with interplanar spacing equal to 14 Å; once the sample was treated with ethylene glycol it produced a strong peak reflection of the smectite and a displacement from \(2\theta = 6.5\) to \(2\theta = 5.4\) (Figure 7). In this case, the peak reflection of the smectite has a maximum intensity of 100% and a maximum spacing of 17 Å. This fact proved that the bentonite of the San José–Los Escullos deposit is expansive.

4.4. Thermogravimetric Analysis

The thermal behaviour of the samples analysed shows similarities and differences, according to the variables in the curves (Figure 8). The curve of sample ZFN-01 shows a higher mass loss compared to the other samples. Like Costafreda [24], the authors have linked this fact to the complex composition of the zeolite, due to its Si/Al ratio, its capacity for cationic exchange and the proportion of smectite–mordenite. The BLT-01 sample is noted because of its mass loss, which is coherent with the data of LOI represented in Table 1. The BFO-01, B-02 and B-03 show similar behaviour which may be related to its low ratio of Si/Al, its high values of LOI and the appreciable content of MgO. The samples B-01 and B-04 behave similarly with regards to their chemical composition, high contents of SiO\(_2\) and Na\(_2\)O and the lowest values of LOI.

Figure 8. TGA thermograms of bentonites from the research and surrounding areas.

4.5. X-ray Fluorescence

The results that stem from the analyses indicate a variable composition in the analysed samples (Table 1). Some bentonites are more siliceous (61.3–64.31% SiO\(_2\)) (e.g., B-01, B-04 and B-05) regarding the less siliceous (51.29–52.53% SiO\(_2\)) (e.g., B-02, B-03). The high SiO\(_2\) contents indicate a marked presence of mordenite in paragenesis with smectite which concurs with what was established by Costafeda [24]; this fact has been proven by comparing the ZFN-01 sample (67.04% SiO\(_2\)), from the San José–Los Escullos deposit, with BLT-01 (47.26% SiO\(_2\)), from the Los Trancos deposit. Garcia-Romero et al. [33] calculated the SiO\(_2\) (51.25–53.47%) content in the Los Trancos bentonites, and found it to be slightly higher than the values obtained in this research; however, their calculations are similar for the Al\(_2\)O\(_3\) (17.01–17.75%), the MgO (4.56–4.87%), the Na\(_2\)O (0.51–1.07%), the K\(_2\)O (0.14–0.43%), the CaO (1.05–1.32%) and the Fe\(_2\)O\(_3\) (2.21–2.41%) (Table 2). The CaO content in the samples from the San José–Los Escullos deposit is discrete and homogeneous with respect to Na\(_2\)O and K\(_2\)O (Figure 9a), emphasising its alkaline character. This conclusion coincides with what was established by predecessor researchers [32,35–37].

Table 2. Chemical composition (wt% oxides) of bentonites from the Los Trancos deposit, by Garcia-Romero et al. [33].

| Sample | SiO\(_2\) | Al\(_2\)O\(_3\) | CaO | Na\(_2\)O | K\(_2\)O | MgO | Fe\(_2\)O\(_3\) | TiO\(_2\) | LOI * |
|--------|---------|---------------|-----|----------|--------|-----|----------------|---------|-------|
| LTBB   | 52.12   | 17.75         | 1.29| 0.51     | 0.14   | 4.56| 2.41          | 0.20    | 20.77 |
| LTBV   | 51.25   | 17.01         | 1.32| 0.51     | 0.43   | 4.78| 2.31          | 0.17    | 20.61 |
| LTBN   | 53.47   | 17.21         | 1.05| 1.07     | 0.27   | 4.87| 2.21          | 0.16    | 19.42 |

*LOI: Loss on ignition.
Figure 9. (a–c). Behaviour of alkaline, alkaline earth and other compounds in bentonites from the San José–Los Escullos deposit and surrounding areas. * Loss on ignition.

The hydrothermal processes displaced the iron ions and concentrated them in the bentonites, producing a marked alteration in the bentonites with typical beige and yellow colours of samples B-01 to 05 and BFO-1 (Figure 3), which differentiates them notably from sample BLT-01. The contents of Fe$_2$O$_3$ (1.3–2.75%), as seen in Table 1, focus of this type of variation in the southern and western sides of the San José–Los Escullos deposit. The mobilisation of the iron ions could have been helped by the placement of subvolcanic bodies of dacitic composition in the peripheries of the Los Frailes caldera [24]; this reasoning fits well with the description made by Martinez et al. [34] and Perez del Villar et al. [38] regarding a dome sited in the Cala del Tomate. These researchers applied the name rubefaction to refer to the ferruginous alteration that some bentonites possess in the south of the Iberian Peninsula.

The MgO contents are even more of an anomaly (4.0–7.67%) in samples B-01 to B-05 (Table 1 and Figure 9b), possibly provoked by a combined process of removal–precipitation of the Mg$^{2+}$ ions from the amphiboles and pyroxene of the dacites and andesites altered by hydrothermal processes.

There is an inverse relationship among SiO$_2$, loss on ignition (LOI) and Si/Al ratio in the analysed samples (Figure 9c). It was observed that the bentonites richer in SiO$_2$ have
lower LOI values and high Si/Al ratios (e.g., B-01, B-04, B-05 and ZFN-01). A different case occurred with the other samples analysed (B-02, B-03, BFO-01, BLT-01) where low SiO₂ contents are related to high LOI values and a low Si/Al ratio. The first case characterises bentonites that are not at all expansive or only slightly, while the second case indicates that the bentonites are expansive. All of the above can be summarised as follows: more siliceous bentonites, with higher Si/Al ratios, have lower loss on ignition (LOI) caused by excess silica (Figure 9c); however, when the Si/Al ratio is low, the samples have a high loss on ignition. Therefore, an inverse ratio was observed, as shown in Figure 9c.

4.6. Scanning Electron Microscopy

Bentonite sample analysis by way of scanning electron microscopy outlined the presence of various types of minerals. These were: smectite, mordenite, halloysite and pyroxene (Figure 10a–d).

Figure 10. Scanning electron microscopy (SEM) micrographs of the bentonite samples (a–d).

Smectite in its montmorillonite variety is the most common mineral, as observed in Figure 10a–d. A comparison between the different samples analysed allowed to establish different morphological properties of the mineral phases. For example, in sample B-01 smectite forms small irregular white to light grey masses (Figure 10a). In sample B-05 it forms light grey and white and can be translucent with compact efflorescence (Figure 10b). In sample BLT-01 it appears to form large compact and irregular masses (Figure 10c), and is practically monomineral, which indicates a high purity. In sample B-02 (Figure 10d) it forms compact aggregates together with mordenite. Both smectite and mordenite developed at the expense of relict minerals such as pyroxene and plagioclase, as well as volcanic glass.
The mordenite found in samples B-05 and B-02 forms elongated crystals >36 μm in size, as well as compact crystalline aggregates (Figure 10b,d). Mordenite is syngenetic with smectite. In Figure 10d both mordenite and smectite grow at the expense of an altered pyroxene crystal.

4.7. Pozzolanicity Test

According to Figure 11a, all the samples analysed have pozzolanic behaviour at 8 days of testing, as their position is below the solubility isotherm (40 °C), as indicated by UNE-EN 196-5:2011 [29]. Specifically, the zeolite sample (ZFN-01) is the most reactive, followed by B-01 and B-05. The other samples, according to their reactivity with Ca(OH)₂, are in the following order: B-04, BFO-01, B-02, B-03 and BLT-01. These data reflect the capacity of the samples to trap the free lime in the solution and the formation of stable reaction products, such as portlandite and tobermorite. The pozzolanic activity is reflected in the decrease of Ca(OH)₂ concentration values from 7.9 mmol/L to 4.1 mmol/L, where sample ZFN-01 is the most pozzolanic. Taking this into account, it should be noted that the bentonite samples contribute effectively to the reaction process. The concentration of the bentonite samples under the curve indicates that B-02, B-03, B-04 and BFO-01 behave similarly, which differentiates them from BLT-01, B-01 and B-05. On the other hand, B-01 and B-05 show behaviour that is quite similar.

According to the data provided in Table 1, B-01 and B-05 are the samples with the highest SiO₂ and Al₂O₃ contents, which is an indispensable condition for a sample to be considered a pozzolan [24]; this is because both SiO₂ and Al₂O₃ represent the acid phase that causes the pozzolanic reaction in a very alkaline solution [29]. Additionally, the high Na₂O and K₂O contents of these samples also confirm the outstanding pozzolanic properties of both samples. Because of this fact, it shows why ZFN-01 is the most pozzolanic reference sample, from which the hierarchy of the quality of each sample studied has been traced.

In addition, it can be deduced that the bentonites from the San José–Los Escullos deposit show a more efficient pozzolanic behaviour than the sample from the Los Trancos deposit (BLT-01) in the 8 days of testing (Figure 11a).

According to the graph in Figure 11b, for 15 days, the pozzolanic reaction of all samples visibly increased. Sample ZFN-01 is still more pozzolanic, followed by B-05, B-01, B-04, BFO-01, B-03, BLT-01 and B-02. The pozzolanic behaviour of BLT-01 stands out in relation to B-02 and B-03 from 8 to 15 days of testing. The set of samples analysed has caused a decrease in Ca(OH)₂ concentration from 4.25 mmol/L to <2 mmol/L at 15 days of analysis.

Figure 11. Graphs of Ca(OH)₂ concentration vs. hydroxyl ions (mmol/L) showing the pozzolanic reactivity at 8 and 15 days (a,b).
Another important aspect to highlight in Figure 11b is the linear trend in the behaviour of the samples analysed, which is different from that observed in Figure 11a. This is interpreted as a consequence of the complex mineralogical, petrological and chemical constitution of the samples discussed in Section 4.1 to Section 4.6, which causes different behaviours in the first stages of reaction with Ca(OH)$_2$. One of the consequences of this is the slow initial reactivity. However, the reactivity becomes stronger with time (15 days), when all components of the samples start to react with Ca(OH)$_2$ (Figure 11b). This aspect is important as these are the same mechanisms that provide mechanical strength to cements, mortars and concretes [24].

The remarkable inclination of the trend line towards the abscissa axis (Figure 11b) is highly relevant in this investigation, because it permits the prediction of an uninterrupted development of the pozzolanic reaction over time. This is interpreted as a decisive factor to use the researched bentonites as pozzolans.

The presence of zeolite in almost all bentonite samples, according to the XRD and SEM tests (Figures 6, 7 and 10), is a very important factor contributing to the reinforcement of pozzolanic properties; this is due to several intrinsic factors: porosity, cation exchange capacity, high active surface area and high adsorption capacity.

According to the thermograms in Figure 8, the bentonites studied lose water and cations with increasing temperature, which activates their ion-exchange capacities and their pozzolanic reactivity.

5. Conclusions

The bentonites from the San José–Los Escullos deposit have pozzolanic behaviour. The complex mineralogical, petrological and chemical constitution reinforces its pozzolanic capacity. These bentonites are able to react with Ca(OH)$_2$ for a long time, which allows the effective neutralisation of free lime in the reaction system and the formation of secondary reaction products, such as tobermorite, which prevents further reaction with sulphates and chlorides. The use of these bentonites as pozzolans could have several advantages, for example, sustainable manufacture of pozzolanic cements, mortars and concretes with high durability and no negative impact on the environment. On the other hand, the specific use of these bentonites in the manufacture of clinker could prevent the emission of CO$_2$ into the atmosphere, thus avoiding the greenhouse effect. Furthermore, concretes and mortars manufactured with bentonite could prevent the degradation of structures built in coastal environments and control the formation of pollutant flows. Bentonite has good pozzolanic capacity compared to zeolite. It is therefore concluded that bentonite and zeolite from the San José–Los Escullos deposit could be mined together, which would increase the productivity and sustainability of the mining activity.

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