Potentiality of clay raw materials from northern Morocco in ceramic industry: Tetouan and Meknes areas

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ABSTRACT
This study aims at evaluating the potential suitability of Tetouan and Meknes (central Morocco) clay material as raw materials in various ceramic applications by investigating their textural, chemical, thermal and firing characteristics. Textural properties were identified by specific surface area, cation exchange capacity (CEC) and bulk density (ρs). Chemical and thermal properties were assessed using XRF and TG/DTA techniques, respectively. Firing characteristics at temperatures from 800°C to 1100°C were determined by linear firing shrinkage, loss on weight and water absorption capacity. The Meknes clays are characterised by medium cation exchange capacity (CEC) and specific surface area (SSA) values due to their moderate smectite content. The Tetouan clays have medium to low CEC and medium SSA values. The main oxides in the clayey samples are SiO₂ (35 – 54.3 wt.%), Al₂O₃ (20.6 – 43.9 wt.%), and Fe₂O₃ (9.7 – 22.4 wt.%). The amount of CaO in Meknes clays ranges from 8 to 12 wt.%, whereas CaO is only present in some Tetouan clay (TE4, TE7, TN4 and TN5). A significant densification of ceramic behaviour could be noticed for most of Tetouan clays at firing temperatures above 1000°C. Meknes clays show earlier densification from 800 °C. The chemical, textural and ceramic properties of Tetouan and Meknes clays indicate their suitability as raw materials for the production of structural ceramics. The high amount of Fe₂O₃ in all clays makes them inappropriate in fine ceramics.

KEYWORDS: Clay materials, Ceramic properties, Ceramic suitability, Morocco.
1. Introduction

Throughout the world, clays are the main raw materials exploited in the fabrication of various ceramic products for building construction. Due to inherently complex physical, chemical and mineralogical characteristics, clays have unique properties related to their own natural genesis [1-4]. In most cases, for economic reasons, the ceramics industry relies on clays from nearby deposits. As a consequence, the characterization and quality control of each clay is important for the technical performance of local products [5, 6].

Pure clays do not occur in nature, they contain mixtures of different clay and associated minerals [7-9]. At present, many ceramic tiles are manufactured from mixtures of mineral raw materials, composed essentially of clays and materials such as quartz, feldspar and carbonates. In the fabrication process, the raw materials are mixed in variable proportions taking into account the influence of each component on the properties of the final products [10, 11]. The components that play fundamental roles for optimum processing, and hence performance of the final products, are clay fraction (< 2µm) for plasticity, feldspar for fluxing and silica as filler material [12-14]. The selection of the appropriate raw materials is based on specific criteria, which are related either to the behaviour during the different stages of manufacturing or to the overall chemical composition. The microstructure and properties of any ceramic depends on the characteristics of the raw materials and processing parameters [15] to assure the quality of ceramic products.

Morocco is among the world top 20 producers and consumers of clayey building materials (http://www.lematin.ma/reader-2007/files/lematin/2011/01/30). The industry of ceramic tiles and bricks is most frequent in northern Morocco. Natural clayey materials are abundant in this region. They play an important economic role with a national production of about 45% for building materials (bricks, ceramic tiles and refractories). However, national ceramic production is still insufficient, and to fill this deficit the Moroccan government has to import ceramic products mainly from Spain, Italy and Egypt. The Tetouan and Meknes area have large construction and development projects to meet the population growth in particular for social and luxury housing construction, which is experiencing a significant economic and tourism development. Meknes and Tetouan clays are currently used for the traditional production of small-scale ceramic factories. Meknes clays are used in small traditional pottery factories using local clays. Artisanal and semi-industrial exploitation of these clays, without any prior study, causes various problems during the manufacturing process, such as deformations and breakage of products. The aim of this study is to evaluate the viability of the Tetouan and Meknes clay deposits as an industrial mineral resource by comparing their textural, chemical and thermal properties as well as drying and firing behaviours.

2. Material

The studied clayey raw materials were collected in the Tetouan and Meknes areas. The Tetouan clay deposit is located in north-western Morocco a few kilometres from the town of Tetouan, in the external Rif domain [17] (Fig. 1). A total of nineteen clay samples were collected in this area. There are two categories of samples: 10 clay samples collected from the exploited quarries labelled (TE) and 9 clays sampled from clay deposits, which are not yet exploited, labelled (TN). The Meknes clay deposit is located in north-central Morocco, about 15 km east of the town of Meknes, in the Sais Basin zone situated in the Pre-Rif domain (Fig. 1). Six Miocene yellow sandy clay samples [18] were collected from the quarries of pottery and artisanal ceramic.

A preliminary characterization of those clay materials was done in a previous study [19]. Tetouan clays are characterized by diversified mineralogical assemblages (in particular a variable proportion of clay, quartz and calcite) in contrast with to Meknes clays (high clay content, quartz and calcite). The clay fraction of Tetouan is dominated by illite and kaolinite with variable contributions of chlorite. Meknes clays are illite and kaolinite, associated with smectite. The studied clay materials consist generally of fine particles with medium to high plasticity and low organic matter content [19].

3. Analytical methods

3.1. Textural properties

The specific surface area of the samples was characterized by the analysis of nitrogen adsorption—desorption isotherms, performed at 77 °K. The measurements were performed using a sorptomatic Carlo Erba 1900, controlled by a computer (Industrial Chemistry, Department of Chemistry, ULg). The analysis
of the isotherms was performed according to the methodology of Vallée, Keller [20], which provided
specific surface area (S_{BET}), micropore volume calculated by the Dubinin–Radushkevich equation
(VDUB), and total pore volume calculated from the adsorbed volume at saturation (Vp). The bulk density
(ρs) was obtained by helium pycnometry on the powdered sample, using Micromeritics Accupyc 1330.
Textural analyses were done at the laboratory of Industrial Chemistry (Department of Chemistry, ULg).
Cation Exchange Capacity (CEC) was measured by the Schollenberger method [21]. The samples were
first saturated by ammonium acetate (1N), and then the ammonium ions in the supernatant were
deprotonated into ammonia with sodium hydroxide solution (0.1N). The ammonia content was then
determined by distillation into a known amount of acid and back-titrated by the Kjeldahl method [22].

3.2. Chemical and thermal properties

The chemical composition of selected elements (Si, Al, Fe, Ca, Mn, Mg, Na, K, Ti, P and S) was
measured as oxides on 2 g of dried and homogenized powder of clayey samples using a Bruker S8 Tiger
wavelength-dispersive X-Ray Fluorescence (WD-XRF) spectrometer equipped with a Rh anticathode
(Industrial Chemistry, Department of Chemistry, ULg). Calibration was made using 35
commercially available certified reference materials of similar matrix (sedimentary rocks, river, lake and
marine sediments, sands and soils). The accuracy ranges from 3 to 7% except for S (25%) and P (20%).
Reproducibilities are above 99% except for S (89%) and P (97%). More details about the method and the
calibration can be found in [23]. The same powdered samples were heated to 1000 °C for 2h to determine
the Loss On Ignition (L.O.I).

Differential scanning calorimetry (DSC) and thermogravimetry (TG) were conducted simultaneously
using a NETZSCH STA 409 PC instrument (Industrial Chemistry, Department of Chemistry, ULg).
Samples were heated from room temperature to 800 °C at 10 °C min⁻¹ under atmospheric air [24].

3.3. Ceramic properties

Industrial tests were carried out as part of the ceramic evaluation process. Clay samples were dried at
105°C for 24 hr and ground to a fine powder and then sieved using a mesh-size of 100μm. Each clay
sample was wetted in order to achieve the proper plasticity for modeling. The samples obtained with these
shaping techniques were 4 cm long, 2 cm wide and 2 cm thick. The drying was done in a shaded and
ventilated room. For different drying times, the mass and the value of the length were measured to
calculate the linear shrinkage. The dried samples (24 h at shaded room plus 12 h at 105°C in oven) were
klin-fired at different temperatures (800, 850, 900, 1000, 1050 and 1100°C) over 1 h. The fired
samples were tested for loss on ignition, shrinkage and water absorption capacity after firing. The linear
shrinkage (LS) was determined following the conventional techniques. The water absorption capacity
(WAC) was determined according to standard procedure UNE 67-027 [25], in fired clay pieces after each
cooking cycle [26]. After preliminary measurements at the end of firing and cooling, the specimens of
each batch were kept dry in an oven until their submission to water absorption. Each dry and cool
specimen was weighed (P1) and the three were then immersed into clean water at 25°C for 24 h. The
specimens were removed from the water, their surfaces were wiped off and the weight (P2) of each was
measured immediately. The water absorption capacity (WAC) was calculated as WAC = (P2 –
P1)/P1×100%.

4. Results and discussion

4.1 Textural properties

Specific surface area (S_{BET}), pore volume (Vp), micropore volume (VDUB) and cation exchange capacity
(CEC) of the samples are listed in Table 1. The surface area of the Meknes samples ranges from 33.3 to
37.9 m² g⁻¹. Sample M1 has the highest specific surface area and M3 the lowest. According to these
specific surface area values, M1 has less illite and smectite content, knowing that the specific surface area
of kaolinite is usually smaller (10-20 m² g⁻¹) with respect to that of illite (80-100 m² g⁻¹) [27], but similar
values for smectite (19 m² g⁻¹) [28]. The Tetouan clays have specific surface area (S_{BET}) values ranging
between 17 and 36 m² g⁻¹. Among them TE1, TE3 and TN9 display the highest SSA values due to their
illite, kaolinite and interstratified clay contents [29].
The CEC values of the Meknes samples (9.0-20.6 meq100⁻¹g) are higher than the Tetouan clays (7.1-18.5 meq100⁻¹g). The high CEC values of the Meknes samples may be explained by their smectite content [19], a mineral characterized by higher CEC value ranging between 80 and 150 meq100⁻¹g [7]. The highest $S_{het}$ and CEC of Meknes and some Tetouan clays (TN and TE) reflect their fine grain size, which is represented by the high amount of clay fraction. The highest $S_{het}$ and CEC of those clays could cause a difficulty to dry and lead to the development of cracks in the drying process. The bulk density ($\rho_s$) of all clay samples are almost similar, ranging from 2.7 to 3.2 gcm⁻³.

4.2 Chemical and thermal properties

The most abundant oxides in the studied samples are SiO₂, Al₂O₃, CaO and Fe₂O₃ (Table 2). According to the major element abundances, the samples are divided into two groups (Fig. 2). The first group is characterized by very low CaO (<1.9%), slightly high Fe₂O₃ (from 12 to 22%) and high Al₂O₃ (from 29 to 44%) in comparison to the second group. On the basis of their low CaO content, the first group is qualified as a non-calcareous clay group, which is mainly represented by the Tetouan clays (TE and TN). The second group displays higher CaO (12-22%), slightly lower Al₂O₃ (20-25%) and lower Fe₂O₃ (8-12%) concentrations and is labelled as a moderate calcareous group. Group 2 consists of all the Meknes samples and TE4, TE7 from the exploited Tetouan samples and TN4 and TN5 from unexploited Tetouan clays. The higher contents of Fe₂O₃ (>5%) in the samples give the reddish color of the clay-based products after firing. The L.O.I at 1000°C (Table 2) ranges from 6 to 24%, which is associated with the presence of clay minerals, hydroxides and organic matter [7, 30]. These material losses are confirmed by the thermal analysis and organic matter content as indicated in Fig. 3 and Table 2.

The DSC/TG (Fig. 3a and 3b) show at least 4 peaks. The first endothermic peak is observed at 90–150 °C for the Tetouan samples and between 96 °C and 113 °C for Meknes samples. Such peaks can be attributed to the removal of adsorbed and interlayered water. The associated mass lost ranges from 0.4 to 0.8% for Meknes clays and between 0.4 and 1.3% for Tetouan clays. A large exothermic peak is observed between 200-450°C. Due to organic matter decomposition [31], it was observed in all studied clays, but was especially well marked at 450°C for M6. A broad endothermic band, sintered at 520 to 550°C for Meknes samples and at 523 to 568°C for Tetouan area samples, is due to clay mineral dehydroxylation and α quartz $\rightarrow$ β quartz transformation [25]. A substantial loss of weight (5–7%) is associated with this endothermic peak for Tetouan samples. As an exception, some Tetouan samples (TN4, TN5, TE7 and TE4) lost less than 3% weight. Meknes samples lost about 4%, demonstrated by elevated clay and quartz content in this area. Small additional endothermic peaks occurred in all samples at 700°C, which are due to carbonate decomposition [32].

4.3. Ceramic properties

a- Drying behaviour

Bigot's curves (Fig.4) were used as preliminary indicators in the choice of raw materials [33] for the ceramic industry. The drying is accompanied by shrinkage of the clay materials, due to porewater loss. The Bigot's curves exhibit the two characteristic phases of the drying process: an initial weight loss with shrinkage and successive weight loss with no further shrinkage. The behaviour of all Meknes clay samples was somewhat similar (Fig.4), although there were some small differences in shrinkage, ranging from 12 to 17%, accompanied by a loss of weight from 19 to 22%. Tetouan clay samples showed a low drying shrinkage (2-6%), in contrast to the Meknes clay samples. Tetouan clays showed a greater increase of the loss on weight (20–23%). The highest linear shrinkage of Meknes clays can be explained by its smectite content and its high PI value [34]. Consequently, Meknes clays have more problematic drying behaviour. By contrast, Tetouan (TN and TE clays) showed a low drying shrinkage, and they have more suitable behaviour.

b- Firing behaviour

The results for linear firing shrinkage, loss on mass and water absorption of the fired clays are displayed in Table 3 and Fig. 5, respectively. Linear firing shrinkage and water absorption have been frequently used as quality and process control parameters in the development and manufacturing stages of the production of structural ceramics such as floor and wall tiles [35, 36]. The linear firing shrinkage could be
used as a direct measure of the extent of densification [37]. The linear firing shrinkage increased gradually from 1000 to 1110°C for most of the Tetouan clays (TN and TE). This evolution reflects densification of the fired clays. As an exception, some fired clays (TE5, TE7 and TE8) do not show any shrinkage. At 800°C, Meknes clays shrank from 17 to 29% and their shrinkage increased again gradually from 850 °C to 1110°C. Meknes clays are characterized by an early densification from 800°C. This is due to the formation of a glassy phase, because they contain large amounts of fluxing agents (Na₂O, K₂O and MgO) (Table 2). Loss on weight stability is indicated from 1000°C for most of the sintered clay bodies (Table 3). It can be seen that an increase in the shrinkage with a decrease in water absorption is associated with an increase in firing temperature. This trend is found especially above 1000°C. Shrinkage is related to the formation of high temperature crystalline phases, and the rates of shrinkage are directly related to the development of glassy material. Formation of gases during vitrification can also have a significant effect on the net dimensions of fired clay minerals. The greatest shrinkage above 1000°C is due to a more significant liquid phase formation. During liquid phase formation, the liquid surface tension and capillarity help to bring particles close together and reduce porosity [38].

A significant decrease of the water absorption is observed for Tetouan clays (TE and TN) from 1000 °C (Fig. 5). As an exception, TE4 and TN4 present the highest water absorption capacity (about 18%). The Meknes clays have high water absorption capacity, ranging from 9 to 21%. At 1100 °C, the water absorption of Tetouan clays decreased to 0.4% and 2.6% for TE6 and TN8, respectively, indicating the total vitrification of those clays. For Meknes clays, the water absorption decreased from 6 to 13%, with the exception of M1, which had 20 % (Fig. 5).

4.4. Suitability for ceramics applications

The suitability of raw clay for their use in the manufacture of ceramic products is determined by mineralogy, chemistry and physical properties of the material. These factors will determine the behaviour of the clay during forming, drying and firing with direct influence on the final product [39]. In order to evaluate the suitability of the studied clays for different ceramic products, a ternary diagram from [40] based on XRD results [19] is shown in figure 6a. Two groups can be identified: those with high quartz contents and those that are rich in oxides, carbonates and feldspars. The mineralogical data for the samples (Fig. 6b) also suggest that most of the Meknes clays can be used for vitrified red floor tile making. However, Tetouan clays can be used for different ceramic applications due to their high quartz amount and their low amount of iron oxide. TN7 and TE9 may be used for making clinker products and TE7 for vitrified red floor tiles (Fig. 6c). Some Tetouan samples (TN5, TN9 and TE7) are located close to the porous light-coloured field; they can therefore be used for porous light-coloured wall tile product. The ternary diagram (Fe₂O₃ + CaO + MgO)/Al₂O₃/(Na₂O + K₂O) (Fig. 6c) is used to classify raw clay materials and industrial ceramic bodies [40]. All Meknes clays plot in the field of red ceramics, close to the field for porous light-coloured ceramics. Most of the Tetouan clays were located outside of the red ceramics field, with the exception of TE4, TE7, TN4 and TN5, which are close to the field for porous light-coloured ceramics. The ideal composition for an optimum white body product (SiO₂ = 72 wt%, Al₂O₃ and total oxides = 8 wt%) was estimated by Fiori et al. [40], Wilson [41] and Baccour et al. [42], who stated that clays containing 8% or more of Fe₂O₃ are red-firing clays. In this context, none of the Meknes and Tetouan clays can be used for the production of fine ceramics, due to their high Fe₂O₃ amount (> 8%). Such application would require processing to reduce this iron oxide content. However, Tetouan clays could be considered as raw materials for structural ceramic products.

5. Conclusion

Tetouan and Meknes clay materials are composed mainly of SiO₂, Al₂O₃, and Fe₂O₃. Meknes clays are characterized by a higher CaO content.
The ceramic behaviours of Tetouan and Meknes clays were interpreted by drying shrinkage, linear firing shrinkage, loss on weight and water absorption. Meknes clays are characterized by a high drying shrinkage, and therefore there will be more difficulty to dry the samples. To solve this problem, sand could be added to reduce their plasticity. Tetouan clays show optimal drying shrinkage. All the fired ceramic properties show moderate changes from 800 to 1000°C. However, significant changes are observed above 1000 °C. An increase in the firing shrinkage and associated with a decrease in the water absorption are especially observed from 1100°C.

Meknes clays and most of Tetouan clays (TE1, TE2, TE3, TE6, TE9 and TN8) can be used on flooring production. In addition some exploited Tetouan clays and currently unexploited Tetouan clays are also suitable for making roofing tiles. Meknes clays and most of the Tetouan clays can be used for different ceramic applications due to their high quartz amount and their small iron oxide amount. Only TN7 and TE9 are being used for making clinker products.

This study demonstrates that exploited Tetouan and Meknes raw clay are suitable for the manufacture of structural ceramic products, with or without additives depending on the ceramic type. New clay outcrops such as unexploited Tetouan clays are also suitable. They constitute a potential ceramic raw material for growing Moroccan ceramic tile industry, and are thus not limited to artisanal production.

Acknowledgements

The WD-XRF data acquisition was made possible at Umeå University through a research grant from the Kempe Foundation to Richard Bindler. The authors would like to thank René Pirard (Laboratoire de Génie Chimique, Université de Liège, Belgium) for the Calorimetry (ATD-TG) and specific surface area (SSA) analyses. We would like to thank Anne Iserentant (Université catholique de Louvain) for the CEC analysis. Finally, our thanks go to Dekayir (Université de Marrakech, Morocco) for his help on the field.

References


Table 1. Textural properties of the clay samples.

<table>
<thead>
<tr>
<th>Samples</th>
<th>$S_{\text{BET}}$ (m$^2$g$^{-1}$)</th>
<th>$V_P$ (cm$^3$g$^{-1}$)</th>
<th>$V_DUB$ (cm$^3$g$^{-1}$)</th>
<th>CEC (meq100$^{-1}$ g)</th>
<th>ps (gcm$^{-3}$)</th>
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<tr>
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Table 2. Chemical composition of the clay samples.

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Exploitable Tetouan clays
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Figures

Figure 1. Structural sketch map of the northern occidental part of the Rif Chain and the Northern central part of Morocco (modified after [45])

1 – Foreland basement, 2 – Meseta and Atlas cover series; 3 – Foredeep basins; 4 – Detached Atlasic cover at Prerif front; 5 – Prerif; 6 – Alpujarrides-Sebtide nappes; 7 – Intrarif, Mesorif, Rif nappes; 8 – Sud-Betic Zone; 9 – Maghrebian Flyschs; 10 – Dorsale calcaire units, 11 – Malague-Ghomaride nappes
Figure 2. Ternary plot of: (a) Al$_2$O$_3$-CaO-Na$_2$O+K$_2$O; and (b) Al$_2$O$_3$-CaO-Fe$_2$O$_3$ (all in wt.%) for studied clay samples.
Figure 3a. Schematic representation of the thermal analysis (ATD/TG) of the clay samples

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↑ Removal of adsorbed water (endothermic peak)
↑↑ Evolution of volatiles and oxidation of organic matter (endothermic peak)
← Quartz transformation and dehydroxylation of phyllosilicates (endothermic peak)
⇌ Kaolinite transformation (endothermic peak)
Figure 3b. Some examples of TG/ DSC curves
Figure 4. Bigot’s diagram of the examined clay samples
Figure 5. Influence of firing temperature on the variation of linear shrinkage and loss on weight.
Figure 6a. Ternary diagram: quartz/ carbonates + Fe-oxides + accessories + feldspars/clays minerals for the studied clays, after [42].

Carbonates, Fer-oxides, Accessories, Feldspars

Quartz

Clay minerals

Exploited Tetouan clays (TE)
Unexploited Tetouan clays (TN)
Meknes clays (M)
Figure 6b. Triangular chart: Quartz + feldspars/calcite + dolomite/phyllosilicates for the studied clay materials, after [42]
Figure 6c. Ternary diagram of Fiori et al. [42] for clays studied: Fe₂O₃ + CaO + MgO/Al₂O₃/(Na₂O + K₂O)

Exploited Tetouan clays (TE)
Not exploited Tetouan clays (TN)
Meknes clays (M)

Porous light-coloured
Red bodies
Clinker