Industrial Potentiality of Alluvial Clays Deposits from Cameroon: Influence of Lateritic Clayey Admixture for Fired Bricks Production

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ABSTRACT

The cartography and characterization of an alluvial clay deposit from Ebebda (Central region of Cameroon) were carried out in order to assess its suitability for the production of fired clay bricks. The clayey area investigated is ~ 50,000 m² with an average thickness of the exploitable layer of 2.2 m, suggesting a deposit of about ~2.2 × 10⁵ tonnes of clay. Mineralogy, physico-chemical and thermal analyses as well as firing properties were performed on representative clay samples. Kaolinite and quartz are the major minerals associated to illite. Upon heating to 900 °C - 1200 °C, the linear shrinkage varies from 1.5% to 15%, the water absorption from 1.5% to 24% and the bending strength from 2 to 12 MPa. The admixture of lateritic clays (widely available) at 50%, 60%, 70% and 80% allow to decrease the shrinkage and bending strength, and to increase the water absorption. Overall, properties were satisfying the requirement for fired brick with 70% of laterite in the mixture.

Keywords: Alluvial Clay Deposit; Characterization; Lateritic Clay Admixture; Brick; Cameroon

1. Introduction

Clays are widely used as raw materials in the field of construction, fine ceramics, pharmaceutical industry, oils decoloration, etc. [1,2]. The evaluation of this natural resource has a ripple effect on the economic development of countries. The potentialities of clayey minerals are generally linked to their structure, composition and physical properties [3]. The exploration of new clay deposits in developing countries such as Cameroon could significantly contribute to socio-economic development [4]. Cameroon’s basement contains industrial clays of various origins (sedimentary, residual and alluvial) [5,6]. Most of previous studies remained preliminary, providing no information on the physico-chemical properties of clays as well as their quantification. This study presents field data of Ebebda’s clay deposit along the Sanaga River. The clay deposit was assessed and the physicochemical and mineralogical compositions were determined in order to evaluate their suitability for the production of fired clay bricks. Laterite residual soils available in Cameroon on nearly 67% of the national territory have been widely studied. Their composition contains Si, Al and Fe with kandites (kaolinite, halloysite) as the main clays minerals. Quartz, goethite and hematite are predominant minerals with rutile and anatase as accessory minerals. All these minerals derive from the weathering of the gneissic basement of the Yaounde’s area [7]. The infrared spectra of kaolinite mineral in lateritic soil present a weak stretching band at 3668 cm⁻¹ reflecting their low crystallinity [8]. The residual clayey layer (upper part of the profile) is common in the Yaounde area [9,10] and according to their typology, morphology and distribution, it is generally observed in the western part of Yaounde on the hills with altitudes between 780 and 800 m. The thickness varies from 3 to 6 m [8]. Despite a few report on the valorization of the laterite clayey soils in low fired brick production (975°C) with no admixture [11], many studies show that the fusibility of these residual clays...
clays and the durability of fired bricks derived cannot be complete due to the mineralogical content of these raw materials [5,8]. In order to better valorize lateritic clayey soil in brick manufacture, the mixture with alluvial clays could present better physical properties. Here, our aim is to test this approach on clayey deposit from central Cameroon. The admixture with lateritic clayey soil was also made in order to reduce the potential production costs. The ceramic properties of the pure fired clay as well as those admixed with lateritic clay were determined.

2. Material and Method

2.1. Material and Geological Outline

The studied alluvial clays are collected from the locality of Ebebda in the central region of Cameroon. The details on studied area are presented on Figure 1. The lateritic clayey soil used as admixture is collected from Etoug-Ebe (3° 54' N, 11° 31' E), a suburb of Yaoundé. The Cameroon’s geology is mainly composed (about 90%) of Precambrian formations dominated by the Ntem Complex and the Panafrican Chain [12,13]. The Sanaga Basin belongs to the Panafrican Chain (500 - 600 Ma) with the outcropping rocks are made of micaschists, gneisses, migmatites and granites known as “Yaoundé series”. Studied clays probably derive from the weathering of basement rocks. These rocks are derived from a high-pressure metamorphism of volcano-sedimentary materials. The weathering products have been undergone erosion, transport and sedimentation along the Sanaga River.

2.2. Method

2.2.1. Clay Deposit Estimation and Sampling

Field campaigns were allowed to establish a geological diagnosis (nature, size, thickness, geometry, homogeneity, accessibility, etc.) of the alluvial clays in the studied area. The exploration and description of outcrops and the cartography of the deposits were done using classical methods of geological survey linked to quantitative evaluation of geomaterials as clays (pit digging, auger drilling, boring holes, etc.). The reserve was estimated on the basis of the drill depth and the studied area [14,15]. During fieldworks, about sixty drillings were performed using a hand auger down to a maximum depth of 5 m. The different steps include: 1) implementation of the sample survey plan, 2) opening of transects providing access to sampling points, 3) realization of 60 pits of 3 to 5 m depth, 4) realization of two observational pits of 1 m² and 2 m depth.

The geographical coordinates of polls and benchmarks were collected using a Garmin type GPS. These coordinates once converted were introduced into the MapInfo software to produce maps. The satellite imagery (Google Earth) was used to georeferencing and to digitize the documents. The mesh used was 50 m in order to produce a map at 1/5000 (Figure 1). The color of the different profile layers was determined using the Munsell code [16]. The samples were selected according to clay facies variation. The “true” density of clay sample was obtained using an automatic pycnometer with helium (type Micromeritics brand AccuPyc, 1330 V 0.3N Norcross USA).

2.2.2. Clay Preparation and Characterization

Average clay samples of selected profile (P2, E2, E6, E14, E18, E46, E50 and E59) and each of different layers of pit (P2a, P2b and P2c) have undergone mineralogical analysis by X-ray diffraction. The sample P2 representing the average of the clay facies (P2a, P2b and P2c) from the pit was also characterized by IR, DTA/DTG, chemical, physical and technological analyses.

X-ray diffraction was performed with a Brucker D8-Advance diffractometer device (copper Kα1 radiation, λ = 1.541838 Å, V = 40 kV, I = 30 mA) at the University of Liege, Belgium.

The measurements were carried out in the 2θ range from 2° to 45° with step size of 0.02° and time per step 2s. Bulk mineralogy and clay fraction were obtained according to [17]. Interpretation of mineral phases was carried out by using Eva software. Qualitative and semi-quantitative estimation are estimated with corrective factors [18-21].

The infrared spectra (IR) were recorded on a Nicolet NEXUS spectrometer. Two milligrams of sample was mixed with 180 mg of KBr to form a tablet used for analysis. Chemical analysis was performed by X-ray Fluorescence Spectroscopy (XRF) with S4 PIONNER Bruker equipment on fine (< 250 µm) powder pellet at Mi-promalo, Cameroun. Simultaneous DTA-DTG experiments were performed using a multidetector high temperature calorimeter (Setaram 85), in dry air condition with a heating rate of 5°C/min from room temperature to 1200°C at ENSCI, Limoges, France.

The grain size distribution was determined by wet sieving of the fraction > 80 µm and gravity sedimentation for the fraction between 80 and 2 µm and the Atterberg limits were obtained by the Casagrande method using ASTM norms, respectively D-422 and D-4318 [22, 23].

Five compositions of bricks were prepared by adding 0%, 50%, 60%, 70% and 80% of laterite (L) in the alluvial clay (P2). The alluvial clay and the laterite were first dried and sieved (Ø < 1 mm) before mixing and hand homogenization. The water content is about 15%, varied from 14% to 15%. The test specimens (80 mm × 40 mm × 18 mm) were made using a 10 KN hydraulic press, then dried at room temperature for 72 h, oven-dried (105°C). The dried samples were fired at 900°C, 1000°C,
Figure 1. Location of Ebebda’s area and sample survey plan for clay deposit along the Sanaga River.
1100°C and 1200°C for a period of 5 hours at a heating rate of 5°C/min in a MultiMate furnace. The firing shrinkage 100(L_{m}−L_{f})/L_{m} were obtained according to the relative variation in the length of the specimen, where L_{m} is the length of the mould and L_{f} is the length of the fired specimen. The water absorption and bending strength were determined using ASTM norm C373-72 and C674-77 respectively [24,25].

3. Results and Discussion

3.1. Field Data and Reserves Estimation of the Alluvial Clay

The elevation of Sanaga valley at Ebebda varied between 200 m and 500 m. The vegetation mainly consists of swamp forest on the right bank while the left bank is mostly covered with savanna. The studies were carried out on both banks. However, the right bank is characterized by a stony soil poor in clay minerals, limiting prospect through depth. The left bank area shows some clay deposits and the profiles are made by five layers (Figure 2): organic soil (0 - 40 cm), fine gray clay mottled with yellow (40 - 130 cm), fine gray clay mottled with red (130 - 180 cm), fine gray clay mottled with light brown (180 - 280 cm) and clayey sand (> 280 cm). For each profile, the thickness of the exploitable layer is from 2 to 2.5 m. The estimated average thickness of clays is ~2.2 m with an approximate area of 5 ha. This gives an average volume of clay per hectare of 22,000 m$^3$. Considering the density of clay (2520 kg/m$^3$) determined by the small endotherm peak at 572°C in a MultiMate furnace. The firing shrinkage of 14 years.

3.2. Mineralogical and Physico-Chemical Characterization

The air-dried XRD pattern of the clays studied (Figure 3) indicates the predominance of quartz (2θ, 27.3°) and kaolinite (2θ, 12.06°). The proportion of the different minerals varies from one sample to another: quartz varies from 43% (P2) to 62% (P2a), K-feldspar from 0% (E50) to 4% (E46), kaolinite from 33% (E59) to 41% (P2c), illite from 5% (E50) to 10% (P2c), smectite from 0% (E2) to 3% (P2), goethite from 1% (E59) to 4% (E2) and hematite from 0% (E2) to 4% (E59). The average data of the main minerals are quartz (49%), kaolinite (33%), illite (7%) and K-feldspar (2%). Goethite and hematite are significantly present (~ 3%) while smectite and anatase are in trace (< 2%) (Table 1).

The IR spectra indicate the presence of two minerals: kaolinite and an oxyhydroxide-rich mineral (Figure 4). The characteristics peaks of kaolinite appear at 3695 cm$^{-1}$ (υ1), 3655 cm$^{-1}$ (υ3) and 3619 cm$^{-1}$ (υ4). A well crystallized kaolinite generally presents four OH-stretching bands: υ1 at 3695 - 3696 cm$^{-1}$; υ2 at 3675 - 3668 cm$^{-1}$; υ3 at 3655 - 3652 cm$^{-1}$ and υ4 at 3619 - 3620 cm$^{-1}$ [26]. The absence of a well defined peak of υ2 (3675 - 3668 cm$^{-1}$) suggests that the kaolinite presents in the alluvial clay (P2) is disordered or poorly crystallized along the (a, b) axes [27]. The disordered or poorly crystallized character of the kaolinite present in the alluvial clay is also evidenced by the absence of Al-OH bending bands at 938 cm$^{-1}$, suggesting stacking faults along the “c” axis [28]. This confirms their sedimentary origin [28,29]. The band at 3434 cm$^{-1}$ is attributed to the OH vibration of hydroxyl alumina [30]. The wide band of low absorption observed around 1626 cm$^{-1}$ is attributed to the presence of water molecules or possibly others radicals such as amorphous constituents of aluminosilicates. The shoulder observed around 2300 cm$^{-1}$ is due to a default of the analytical device.

The thermal curves (DTG/DTA) of the alluvial clay (Figure 5) have evidenced between 0°C - 200°C, a first weight loss (about 2%) due to the departure of the hygroscopic water also materialized by the first small endotherm peak at 72°C. The small endotherm peak observed at 249°C with 0.78% of weight loss could arise from the dehydration of iron or alumina oxhydroxides. Between 400°C and 600°C, there is a continuous weight loss (7.22%) due to the departure of the structural water of kaolinite (second endothermic at 493°C). This particular weight loss is generally linked to the amount of kaolinite present in the clay [31]. On the other hand, the broad exothermic band between 100°C and 300°C is linked to the combustion of organic matter and the small endotherm peak at 572°C is due to the trans-

![Figure 2. Profile of alluvial clay at the locality of Ebebda, Cameroon.](image-url)
formation of quartz alpha to quartz beta [32]. The LOI result confirms the presence of organic matter which represents the volatile materials. No further weight loss is observed above 800°C. The last small endothermic peak observed between 800°C and 1000°C is due to the production of silicate liquid deriving from the melting of feldspar, while the exothermic band corresponds to the formation of spinel or mullite phase [33].

The physico-chemical properties of the alluvial clay and the laterite used for the technological tests (Table 2) show SiO₂ (60%), Al₂O₃ (14%) and Fe₂O₃ (7%) as the predominant oxides of the alluvial clay. The lateritic soil used contains also the same oxides, but it is less siliceous (40%) and is richer in alumina (32%) and iron (12%). The sum of bases (K₂O, MgO, CaO) is low in the lateritic soil (< 1%), but slightly higher (1.37%) in the alluvial clay.

The particle size distribution of the alluvial clay is

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**Figure 3.** Bulk mineralogy of alluvial clays from the locality of Ebebda, Cameroon.

**Table 1.** Some properties of used clayey soils.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Depth (cm)</th>
<th>Total Clay</th>
<th>Quartz</th>
<th>K-Feldspath</th>
<th>Anatase</th>
<th>Goethite</th>
<th>Hématite</th>
<th>Kaolinite</th>
<th>Illite</th>
<th>Smectite</th>
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<tbody>
<tr>
<td>P2</td>
<td>30 - 300</td>
<td>46</td>
<td>43</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>36</td>
<td>9</td>
<td>3</td>
</tr>
<tr>
<td>P2a</td>
<td>30 - 100</td>
<td>30</td>
<td>62</td>
<td>3</td>
<td>1</td>
<td>3</td>
<td>0</td>
<td>24</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>P2b</td>
<td>100 - 230</td>
<td>41</td>
<td>48</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>4</td>
<td>34</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>P2c</td>
<td>230 - 300</td>
<td>53</td>
<td>43</td>
<td>0</td>
<td>2</td>
<td>2</td>
<td>0</td>
<td>41</td>
<td>10</td>
<td>2</td>
</tr>
<tr>
<td>E2</td>
<td>40 - 180</td>
<td>43</td>
<td>49</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>4</td>
<td>35</td>
<td>7</td>
<td>0</td>
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<td>150 - 290</td>
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<td>7</td>
<td>2</td>
</tr>
<tr>
<td>E14</td>
<td>15 - 100</td>
<td>43</td>
<td>49</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>33</td>
<td>8</td>
<td>1</td>
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<tr>
<td>E18</td>
<td>90 - 220</td>
<td>42</td>
<td>48</td>
<td>2</td>
<td>2</td>
<td>4</td>
<td>3</td>
<td>33</td>
<td>8</td>
<td>0</td>
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<tr>
<td>E46</td>
<td>80 - 160</td>
<td>44</td>
<td>47</td>
<td>4</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>35</td>
<td>7</td>
<td>2</td>
</tr>
<tr>
<td>E50</td>
<td>100 - 170</td>
<td>42</td>
<td>51</td>
<td>0</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>36</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td>E59</td>
<td>170 - 190</td>
<td>34</td>
<td>51</td>
<td>4</td>
<td>3</td>
<td>1</td>
<td>4</td>
<td>24</td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td>Average</td>
<td>42</td>
<td>49</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>33</td>
<td>7</td>
<td>1</td>
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Table 2. Properties of studied clays.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Density g/cm³</th>
<th>Clay</th>
<th>Silt</th>
<th>Sand</th>
<th>Fines</th>
<th>WI</th>
<th>Wp</th>
<th>Ip</th>
<th>SiO₂</th>
<th>Al₂O₃</th>
<th>Fe₂O₃</th>
<th>CaO</th>
<th>MgO</th>
<th>K₂O</th>
<th>TiO₂</th>
<th>LOI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alluvial clay (P2)</td>
<td>2.52</td>
<td>72.9</td>
<td>19.6</td>
<td>7.34</td>
<td>89.7</td>
<td>82.1</td>
<td>35.8</td>
<td>46.3</td>
<td>61.5</td>
<td>14.7</td>
<td>7.6</td>
<td>0.05</td>
<td>0.28</td>
<td>1.19</td>
<td>1.38</td>
<td>11.89</td>
</tr>
<tr>
<td>Lateritic soil (L)</td>
<td>-</td>
<td>35.0</td>
<td>23.0</td>
<td>41.0</td>
<td>59.0</td>
<td>41.6</td>
<td>18.7</td>
<td>40.6</td>
<td>32.1</td>
<td>12.0</td>
<td>0.23</td>
<td>0.44</td>
<td>0.17</td>
<td>1.48</td>
<td>11.12</td>
<td></td>
</tr>
</tbody>
</table>

*composition of lateritic clay is for [8].

Figure 4. IR curves of alluvial clay from pit P2.

Figure 5. Thermal curves of alluvial clay from pit P2.

70% of clay fraction, 15% of silt and 10% of sand. The greatest amount of fines particles (90%) suggests that this soil belongs to the very heavy clay group according to the Belgian texture of soil [34]. For the lateritic clay (L), [8] reported 35% of clay fraction, 25% of silt, 40% of sand and trace of gravel (1% - 2%). In the Casagrand diagram (Figure 6), the alluvial clay belongs to the field of plastic clay while the lateritic clayey soil belongs to the medium plastic group. The great proportion (70%) of the clays fractions (< 2 µm) in the alluvial clay associated with their high plasticity (46%) favor a high shrinkage, which make them unsuitable (at the pure state) for bricks manufacture [2]. In the ceramic industry, the plastic clays are generally used to amend non fusible clays [2,8,35]. Since lateritic clay soil do not integrate any fluxing min-

Figure 6. Position of alluvial clay “P2” and lateritic clay soil “L” in the Casagrand diagram.

erals, the presence of feldspar, illite and smectite in the alluvial clay could assure a good fusibility of the bricks resulting from the mixture of the alluvial clay and laterite.

3.3. Technological Properties of Bricks

The linear shrinkage, water absorption and the flexural strength were used to access the fusibility of the mixture formulations. The values of linear shrinkage of the bricks increase with increasing the firing temperature (Figure 7). At 900°C, the values are less than 2%, rising to 3% - 4% at 1000°C and ~ 8% for brick resulting from the composition of 100% alluvial clay (L0%). Beyond 1000°C, the shrinkage is very high (> 8%) for all the mixtures. This shrinking behavior, primarily attributed to the degree of sintering, is probably due to rearrangement reactions within the firing product associated to partial sintering of some phases [36]. The 100% alluvial clay formulation (L0%) has a higher linear shrinkage comparatively to the pure clay admixed with laterite, suggesting a positive effect of laterite addition in the shrinkage.

The water absorption values decrease with increasing temperature (Figure 8). These values are in the range 20% - 25% at 900°C - 1000°C and below 20% above 1000°C. The addition of laterite slightly increases the percentage of water absorption, probably due to the relatively refractory character of the laterite, which mitigates the formation of vitreous phases at higher temperatures.
firing temperature require is 1050°C for industrial production of fired bricks and the minimal temperature for the production of fired bricks. The alluvial clay studied, admixed with the laterite, could be used for the production of fired bricks. The alluvial clay exhibits a bending strength of 2 MPa - 7 MPa, rising to 6 MPa - 12 MPa for temperatures above 1000°C. The admixture of laterite in the alluvial clay slightly decreases the values of bending strength at 900°C - 1100°C, this may arise from the ongoing reaction of the Fe-bearing phases at this temperature [39]. According to Sigg (1991) [40], 7 MPa is the minimum value required for normal fired brick and 1050°C was reported as the maximum temperature for the production of fired bricks. The alluvial clay studied, admixed with the laterite, could be used for industrial production of fired bricks and the minimal firing temperature require is 1050°C.

4. Conclusion

The detailed mapping of Ebebda’s clay site has been carried out, followed by some characterization. Ebebda’s area offers a great potentiality for the exploitation of raw clay materials for the ceramic industry, particularly on the left bank of the Sanaga River. The clayey layer (average thickness ~ 2.2 m) is overlain by a thin layer of top soil. The clay material is estimated at 2.2 × 10⁶ tonnes over the studied area (05 hectares). The mineralogical paragenesis is made by quartz (40%), kaolinite (30%) and illite (7%). K-feldspar, goethite, hematite and smectite are also present. From the physico-chemical properties, the alluvial clays were classified as plastic clays. Our experiments demonstrate that the admixture of lateritic clay has a positive effect in reducing the shrinkage. However, the presence of lateritic clays slightly increases the water absorption of the bricks and decreases the bending strength. The best properties were obtained for a mixture of 70% of laterite with 30% of alluvial clays.

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