Geophagic Clayey Materials of Sabga Locality (North West Cameroon): Genesis and Medical Interest

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Abstract: Geophagia, the deliberate ingestion of clayey materials, is a complex eating behaviour with obscure etiology and numerous health/medical problems. Geological, mineralogical and geochemical studies were carried out on Sabga geophagic clayey materials located within a trachyte Bamenda mountain, about 16 km east of Bamenda town (North West Cameroon), in order to define the genesis, and its medical interest. Four (04) samples were characterized by different techniques: description of outcrops and pits, particle size distribution by laser dispersion, X-ray diffraction, Fourier transform infrared spectrometry, Scanning electron microscopy and bulk chemical analyses. The main clay minerals were smectite (49-60%) and kaolinite (4-6%). The other major minerals are quartz (19-34%), feldspar (6-12%), goethite (3-6%), and hematite (1-3%). The average particle diameter varied from 2 to 9 µm. Results from the different analytical techniques point out a meteoric weathering of feldspar as petrogenetic origin of geophagic clayey materials. Smectite and kaolinite, which are the main desirable minerals components of the geophagic clayey materials, were geochemically derived from surrounding trachytes. However the presence of crystalline silica (quartz) up to 2% limits the application of some Sabga geophagic clayey materials for pharmaceutical clays. The high radioactive (Th, U), carcinogenic (Cr, Cu, Pb, Ni) and teratogenic (or birth defects) (Cu, Zn, Pb) elements contents in some Sabga geophagic clayey materials implied some pretreatment.

Keywords: Geophagic Clayey Materials, Meteoric Weathering, Sabga, Trachyte

1. Introduction

The consumption of clayey materials occurs in almost all the continents in the world including North America [32], Central America [36], South America [1], Asia [2], Europe [35] and Africa [64, 23, 3, 44]. The practice of consuming geophagic clayey materials is very common in many African countries including Cameroon, where several localities abound in edible geophagic clayey materials. Among the different clay minerals, kaolinite in Koussérie, Mayouom, Dschang and Moko; smectite in Baba and Sabga; and halloysite in Balengou [63, 60, 24, 25, 17, 18, 3, 22, 44] have been reported in Cameroon. As far as pharmaceutical applications of Sabga are concerned, we should keep in mind that fine smectite from Sabga is ingested by neighbouring populations (mainly by pregnant women).

Smectite and kaolinite are used in modern pharmaceuticals to prevent nausea, vomiting, and gastrointestinal disorders [96, 18, 3, 44]. Geophagia has been of interest to researchers and analysts due to continued habitual, religious and cultural practices by diverse global communities in spite of reports contraindicating its purported benefits. Most geophagic
clayey materials are highly retentive due to their fine-grained sizes allowing their absorption of metal ions. The concentration levels of these elements determine their usefulness or harmfulness when consumed by humans. Geophagic clayey materials are considered as sources of elemental supplementation especially Fe, Ca, K and Zn, as well as toxic metals such as As and Pb [50].

No known studies have been carried out to characterize the Sabga geophagic clayey materials. Thus, neither the genesis nor the suitability of clays for medicinal applications has been carried out. Effective valorisation of monomineral clays such as kaolin, talc, and smectite as raw materials for applications in the pharmaceutical, ceramic, paint, and paper industries among others often needs knowledge of the clay deposit genesis [58, 65] and the mineral quality [6, 56, 53], and geochemical composition [24]. The mineralogical inertness dictates that the silica polymorphs (quartz) should not exceed 2% [89], because they have been considered responsible for carcinogenicity [38].

A wide range and variety of minerals are used as excipients in pharmaceutical preparations including oxides (rutile, zince, periclase, hematite, maghemite, magnetite), hydroxides (goethite), carbonates (calcite, magnesite), sulphates (gypsum, anhydrite), chlorides (halite, sylvite), phosphates (hydroxyapatite), phyllosilicates (smectites, palygorskite, sepiolite, kaolinite and talc) and recently tectosilicates (zeolites) [13]. Within the smectite group, montmorillonite, saponite, and hectorite are the most widely used clay minerals. Smectites can be used as disintegrant, diluent and binder, emulsifying, thickening and anticaking agent, and carrier-releaser in pharmaceutical preparations [13].

The term “bentonite” is too generic, as it can be used both for a rock consisting mainly of smectites (mineralogy) or a material mainly containing montmorillonite (pharmacy).

The focus of this study was therefore to mineralogically and geochemically analyse geophagic clayey materials from Sabga locality in the physical state in which they are ingested, in order to elucidate on the deposit quality, its genesis and possible medicinal utilisation. It is anticipated that the findings of the study could establish baseline mineralogical and geochemical characterizations of Sabga geophagic clayey materials; and contribute to the renewed and recently rekindled research interests [92], and intellectual debates on geophagic practice within the broader scientific community [29, 77].

2. Geographic and Geological Setting

Sabga area belongs to the Bamenda Highlands and is sited at latitudes 5.58° to 6.03°N and longitudes 10.19° to 10.25°E (Figures 1c and 2a). This locality is located about 16 km east of Bamenda town, North West Cameroon. The relief is hilly with an altitude range between 1180 and 2050 m and separated by U and V valleys. The hydrographic network shows dendritic drainage. The climate is Sudanese tropical, with temperatures between 17 and 23°C; and annual mean rainfall of 2396 mm. This area is also characterised by geothermo and mineral springs [14].

Figure 1. Geological map of the Bamenda Mountains showing the location of study area [42] (a) Location of Cameroon in Africa; (b) location of the Bamenda Mountains along the Cameroon line showing the outcrop of the Cameroon line volcanic rocks (black) (c) Location of the western highlands. C = Cameroon. COA = West African Craton; CC = Congo Craton; CT = Tanzanian Craton; CK = Kalahari Craton. African Cratons.
This climate allows the formation of colluvial deposits soils on magmatic and metamorphic basements around the Sabga Mountain. The soils have been studied for potential oil bleaching earth, adsorbents for mercury (II) species, adsorption of lead (II) ions utilized on a commercial scale to decontaminate industrial effluents [82, 45, 47]. Detailed geological, pedological data have not been reported on the studied area.

The area under study is part of the Bamenda Mountains, situated on the Cameroon Volcanic Line (CVL). Sabga locality is located between the Bamboutos Mountains to the South West and the Oku massive to the North West (Figure 1). This volcanic province is made up of both mafic and felsic rocks lying on a Panafircan or older basement [41]. The felsic lavas are mainly represented by trachytes, with subordinated benmoreites and alkaline to peralkaline rhyolites [40], whereas the mafic rocks are basalts to mugearites [41]. The area is made up of rock pedestals (plutono-metamorphic) with volcanic rocks and alluvium overlying on top of it. The volcanic rocks formations are made up of trachytes, rhyolites, ignimbrites and basalts [43, 14, 26]. Trachytes appear exclusively at the mountain peaks. The expected parent rock of the geophagic clayey materials is a trachyte which is in the form of slabs and is greenish, massive and encloses laths of feldspar oriented in the same direction as the typical trachytic structure. The rock displays a microporphryric texture and is composed of pyroxens, feldspar and plagioclase [43]. Trachyte is highly fractured thereby facilitating weathering.

### 3. Methods

#### 3.1. Field Observation of the Geophagic Clayey Materials

![Figure 2](#) Detail geological map of Sabga geophagic clayey materials (with the location of analysed samples) and related interpretative sketch cross section (A-A').
The gangue constituted by organo-mineral and lateritic soil layers. A detail geological map after the field mapping is shown on Figure 2a. Cross section illustrates the structure of the geophagic clayey materials (Figure 2b). The whole soil section shows 5 successive layers. From the top to bottom, they include: An brown organic matter-rich A horizon (0-0.1 m) with a sandy loam texture and crumb structure; a thick B horizon made up of a lateritic surface and mottled subsurface composed of a 0.1-0.5 m thick, reddish brown (7.5R4/6), and reddish yellow (7.5 YR 6/8), with clayey sand, gravels and abundant quartz grains. The transition to the saprolite below is not gradual. A weathered C horizon is constituted of three layers, with very thick (0.5-3.5 m). Saprolite is represented by these three layers. The first layer is grey whitish (2.5Y 8/1), compact with a 0.5-2 m thick and clayey texture. The second layer is yellow (10YR 7/8) mixed with grey and white patches, with clayey texture and a 2-2.5 m thick. The third layer is whitish grey (2.5Y 8/1), very compact with clayey sand texture and a 2.5-3.5 m thick, spotted with yellow nodules with many slightly weathered fragments of trachyte. At mid-way of the Sabga hill formed by trachyte, the geophagic clayey materials outcrops in small interfluves, thus constituting nest of ore easily identified by the white, grey yellow and red soil. They outcrop on the Sabga hill some eight km from different escarpments. The representative vertical cross section presents from the top to the base (Figure 3).

3.2. Sampling and Samples

On the Sabga clay deposit mine, four pits of varying depths of 4 to 10 m were dug. Samples of geophagic clayey materials commonly ingested by individuals in Sabga locality were obtained from different sites. The samples were the true representatives of the different geophagic clayey materials available in Sabga locality. Four geophagic clayey material samples (SA, SB, SC, SD) were collected for laboratory analyses (Figure 2).

3.3. Laboratory Analyses

Laboratory analyses conducted included particle size distribution by laser dispersion, X-ray diffraction (XRD), Fourier transform infrared (FTIR) spectroscopy, scanning electron microscopy (SEM) and major and trace elements

![Image](https://via.placeholder.com/150)
compositions analyses.

Particle size distribution of the geophagic clayey materials was determined by laser light scattering using a Malvern Masterizer 2000 particle size analyzer at the Laboratory of "Chimie" University of Liège in Belgium. A suspension of each sample was then loaded into a Malvern Mastersizer 2000 fitted with a Hydro 2000G dispersion unit. A polydisperse mode of analysis and a refractive index of 1.53 with an adsorption of 0.1 were chosen. Size data collection was performed at constant obscuration in the range 10–20%. Scattered light data were recorded from 2000 to 5000 snapshots of 10 µs. The correlation between the angles of light scattered from the particles in a laser beam was used to determine the size distribution of these particles. Particle data generated for each soil sample included the PSD and derived diameters D10, D50 and D90, where D10, D50 and D90 represent the 10th percentile, median diameter and 90th percentiles respectively of the geophagic clayey material sample. Values for D10 and D90 describe the maximum particle diameter below which 10% and 90% respectively of each sample volume exist.

Mineralogical analysis were carried out at the Laboratory of "Argiles, Géochimie et Environments Sédimentaires", University of Liege (AGEs, Belgium). X-ray diffraction patterns were done with a Bruker Advance D8 diffractometer using a Cu- Kα radiation under 40 kV and 30 mA operating conditions. Bulk powder was studied following normal procedure of clay analysis as presented in [57]. The mineral percentages in different samples are calculated using the height of a diagnostic peak multiplied by a corrective factor [12, 8] together with the so-called 100% approach (the term 100% approach connotes that the sum of all phase quantities identified in a sample is 100%). The common reflection of total clays at 4.47 Å is multiplied by a corrective factor of 20 to give an estimate of the total clays in the bulk sample. For the semi-quantification of the different clay minerals and non-clay minerals, the estimation of smectite and kaolinite were based on the height of 001 reflections (at ~14 and ~7 Å, respectively) with a correction factor of 0.7 and 0.25, respectively [27] on glycolated specimens estimation of quartz, feldspar, goethite and hematite were based on the height of 001 reflections (3.35, 3.24, 4.97 and 2.07 Å respectively) with a correction factor of 1, 4.3, 7 and ~3.33 respectively [12, 8]. For clay fractions, three patterns were recorded in sequence under natural condition, after saturation with ethylene-glycol for 22 to 24 h, and heating at 500°C for 4 h. Mineral identification was processed using EVA software. Semi-quantitative mineralogical composition of bulk powder was estimated from the intensity of a diagnostic peak multiplied by a corrective factor [12, 8]. For the clay fraction, semi quantitative estimation was based on the height of specific reflections, generally measured on EG runs [5]. Smectite in natural condition appearing at 14 Å and in the glycolated preparation, gives a very strong (001) reflection at 17 Å [57]. After heating, the smectite peak collapses at 10 Å (Figure 5a, b). Kaolinite is identified by the disappearance spacing around 7 Å on heating (Figure 5b).

Fourier transform infrared spectra were recorded in the 4000–4000 cm\(^{-1}\) range on Bruker FTIR Spectrometer at the Laboratory of "Minéralogie" University of Liège in Belgium, using pressed pellets heated to 110°C overnight. The disks of 2 cm diameter were prepared by mixing 10 mg sample with 150 mg KBr.

The scanning electron microscopy (SEM) was used to observe their morphology and size; the observations were carried out on an ESEM Philips XL-30 device at laboratory of "Chimie Inorganique Structurale" (University of Liege). The images were obtained by a secondary electron detector after metallization with gold powder by plasma spraying for an accelerating voltage scan.

Major elements analyses were performed at the laboratory of "Pétrologie Sédimentaire (PETROSED)" at the University of Liège in Belgium by emission spectrometry using inductively coupled plasma and atomic emission source (ICP–AES). Trace elements and rare earths analyses were performed at the laboratory of Géochimie Environnement de Surface Gestion des Collections, Musée Royale d’Afrique Centrale Bruxelles in Belgium by mass spectrometry (ICP–MS) after fusion with LiBO\(_2\) and dissolution in HNO\(_3\).

4. Results

4.1. Qualitative Mineralogy of Geophagic Clayey Materials

The main minerals identified by XRD in geophagic clayey materials (Figure 4) are a 14.12 Å phase considered as 2:1 phyllosilicate group mineral, a 7.18 Å phase assigned to kaolinite and characteristic peaks of feldspar and quartz. The 14.12 Å phase is the predominant mineral in all the samples. Goethite and hematite had weak or few characteristic peaks in some samples.

The SEM micrographs show that geophagic clayey materials are characterized by the presence of pseudo-spherical smectite aggregates with different shapes and sizes (Figure 6a, b, d). The smectite particles forming the aggregates are less than 10 µm in size and they are characterized by corn-flake morphologies (Figure 6b, c).

The FTIR spectra of geophagic clayey materials samples (Figure 7) show the OH-stretching vibrations region (3700–3400 cm\(^{-1}\)). The infrared absorption spectra show a sharp band around 3620 cm\(^{-1}\) due to OH stretching (Figure 7). This band, clearly visible on Figure 7b, c, d, corresponds to AIAOH stretching of smectite [48]. The band at 3697 cm\(^{-1}\) (Figure 7b, d) are characteristic of kaolinite [93], and 3620 cm\(^{-1}\) is related to the presence of montmorillonite [49]. The Si-O stretching band is observed in the IR spectra of both di- and tri-octahedral smectites, montmorillonite at 1030 cm\(^{-1}\)-1035 cm\(^{-1}\). The presence of quartz in all samples is confirmed by Si–O symmetrical stretching vibrations around 792 cm\(^{-1}\) and 795 cm\(^{-1}\). The bands at 1637 cm\(^{-1}\)-1641 cm\(^{-1}\) are due to adsorbed water [33].
Figure 4. XRD patterns of Bulk powder. Sm = Smectite; Kaol = Kaolinite; TCM = Total Clay Mineral; Fd = Feldspar; Qtz = Quartz; He = Hematite; Go = Goethite.

Figure 5. XRD patterns of clay fraction. Black pattern = untreated; Blue pattern = solvated by ethylene glycol; Red pattern = heat-treated at 500°C; Sm = Smectite; Kaol = Kaolinite.
Figure 6. Some examples of the smectite micromorphology found in Sabga geophagic clayey material samples. a, b) SEM of sample SA; c, d) SEM of sample SB.

Figure 7. FTIR spectra for geophagic clayey materials from Sabga.
4.2. Semi-Quantitative Mineralogy of Geophagic Clayey Materials

Results of mineralogical reconstitution are shown in Table 1. The samples display high smectite and kaolinite contents, 49–60% for smectite and 4–6% for kaolinite. Associated minerals are feldspar 6–12%, quartz (19–34%), goethite 3–6%, and hematite 1–3%.

4.3. Particle Size Distribution of Geophagic Clayey Materials

Values for the median diameter of the particles making up the Sabga geophagic clayey materials ranged from 1.8–9.3 µm, while values of D10 and D90 for of the particles making up the geophagic clayey materials ranged from 0.85–132 µm and 4–60 µm respectively (Figure 8). The derived diameters of individual samples are presented in Table 2.

4.4. Geochemistry of Geophagic Clayey Materials

4.4.1. Major Oxides

All samples had silica content ranging from 41.52–47.89 wt%. These chemical characteristics correlate with quartz content. The alumina (Al2O3) contents in the soils were also slightly higher with a range of 11.03–23.41 wt% (Table 3). Samples also had very high values for the Fe oxides (4.42% in sample SC to 10.62% in sample SD) (Table 3), inducing yellowish and reddish colour of samples.

| Samples | Smectite (%) | Kaolinite (%) | Total Clay Mineral (%) | Quartz (%) | Feldspar (%) | Goethite (%) | Hematite (%) |
|---------|--------------|---------------|------------------------|------------|-------------|-------------|-------------|
| SA      | 60           | 0             | 60                     | 19         | 12          | 6           | 3           |
| SB      | 52           | 6             | 58                     | 32         | 6           | 3           | 1           |
| SC      | 49           | 4             | 53                     | 34         | 10          | 4           | 1           |
| SD      | 59           | 0             | 59                     | 26         | 8           | 4           | 3           |

4.4.2. Trace Elements

For trace elements compositions of studied geophagic clayey materials (Table 3), some trends are present. The concentrations of Y, Nd, Ce, Tb, La, Zn and Zr were high (Table 3). As, Cd, Co, Ge, U and Ni had the lowest concentrations with all values being < 10 ppm (Table 3). The concentrations of other elements (Cu, Ga, Hf, Pb, Sc, Sm, Ta and Th) were between 10 and 75 ppm (Table 3).

Trace elements such as Co, Cu and Zn which are of nutritional significance to humans and Ni, Pb, and Cr which have lower toxicity thresholds were relatively low compared to those found in soils for agricultural purpose [61]. Attention must be drawn to the amounts of Pb and As in the pharmacopeia limits for “Bentonite” and “Purified Bentonite”. Limits for Pb are 40 and 15 ppm, and for As the limits are 5 and 3 ppm respectively [83, 84]. However, trace element analyses (Table 3) show notable amounts of radioactive (Th, U), carcinogenic (Cr, Cu, Pb, Ni) and teratogenic (or birth defects) (Cd, Cu, Zn, Pb) elements, higher than those reported in recommended pharmaceutical clays [51]. The REE content normalized to Sabga trachyte [42] and chondrite CI values of [52] are shown in (Figure 9a, b). The trachyte normalised trace elements patterns for representative geophagic clayey materials from the Sabga locality are parallel and flat with negative Eu and Ce anomalies (Figure 9a). The chondrite normalised trace elements patterns for representative geophagic clayey materials from the Sabga locality show similar REE patterns, being more enriched in LREE compared to HREE with distinct negative Ce and Eu anomalies (Figure 9b).
### Table 3. Major elements and trace elements compositions of geophagic clayey materials from Sabga compare with a pharmaceutical clay composition (sample named Pharm Clay is from [46] and [51]).

| Samples          | SA    | SB    | SC    | SD    | Pharm Clay |
|------------------|-------|-------|-------|-------|------------|
| SiO₂ (wt.%)      | 41.52 | 47.11 | 47.73 | 47.89 | 44.6–46.4  |
| Al₂O₃             | 11.03 | 23.41 | 21.18 | 17.13 | 38.1–39.5  |
| Fe₂O₃             | 4.64  | 7.65  | 4.42  | 10.62 | 0.1–0.2    |
| MnO              | 0.12  | 0.01  | 0.11  | 0.12  | -          |
| MgO              | 0.21  | 0.00  | 0.36  | 0.42  | 0.1–0.2    |
| CaO              | 0.34  | 0.15  | 0.56  | 0.79  | 0.1–0.2    |
| Na₂O             | 0.33  | 0.10  | 0.77  | 0.33  | 0–0.1      |
| K₂O              | 1.22  | 0.75  | 1.56  | 1.42  | 0.2–0.2    |
| TiO₂              | 0.27  | 0.74  | 0.29  | 0.67  | 0.1–1.4    |
| P₂O₅              | 0.07  | 0.05  | 0.03  | 0.03  | -          |
| LOI = Loss of ignition (wt.%) | 15.52 | 19.60 | 20.55 | 20.35 | 13.8–13.9  |
| Total            | 94.59 | 99.20 | 98.55 | 98.97 | -          |
| Al₂O₃/MgO         | 52.52 | 0.00  | 58.8  | 40.7  | -          |
| MgO/Fe₂O₃         | 0.04  | 0.00  | 0.08  | 0.04  | -          |
| Trace elements (ppm) |       |       |       |       |            |
| As (<|dl)      |       |       |       |       | <0.3       |
| Sc              | 11.8  | 10.4  | 13.8  | 12.5  | 10.2       |
| V               | 4.7   | 2.6   | 2.6   | 1.0   | 24         |
| Cr              | 7.0   | 1.9   | 3.1   | 3.5   | 5.4        |
| Co              | 1.3   | 0.9   | 0.8   | 2.1   | 5          |
| Cd (<|dl)      |       |       | <|dl|  <|dl|  0.02     |
| Ni              | 7.5   | 4.2   | 3.1   | 4.3   | 5          |
| Cu              | 18.7  | 23    | 39    | 63    | 4.71       |
| Zn              | 248   | 147   | 177   | 989   | 9          |
| Ga              | 48    | 69    | 42    | 84    | 1.79       |
| Ge              | 4.4   | 3.8   | 1.4   | 3.0   | n.d        |
| Rb              | 82    | 14.5  | 10.0  | 56    | 83         |
| Sr              | 48    | 11.4  | 39    | 55    | 100        |
| Y               | 123   | 137   | 132   | 200   | 29         |
| Zr              | 927   | 1648  | 624   | 3114  | 162        |
| Nb              | 151   | 260   | 101   | 520   | n.d        |
| Cs              | 4.8   | 2.1   | 1.00  | 3.8   | -          |
| Ba              | 76    | 46    | 108   | 98    | 147        |
| Bi (<|dl)      |       |       | <|dl|  <|dl|  0.16     |
| La              | 122   | 249   | 277   | 459   | 25.8       |
| Ce              | 169   | 242   | 120   | 540   | 46         |
| Pr              | 27    | 66    | 67    | 115   | n.d        |
| Nd              | 99    | 241   | 241   | 403   | 19         |
| Sm              | 18.8  | 45    | 41    | 69    | 4.7        |
| Eu              | 1.07  | 4.6   | 2.4   | 3.5   | 0.8        |
| Gd              | 18.7  | 34    | 32    | 49    | n.d        |
| Tb              | 1.72  | 2.19  | 1.56  | 2.08  | 0.7        |
| Dy              | 19.5  | 26    | 24    | 36    | n.d        |
| Ho              | 3.9   | 4.6   | 4.3   | 6.3   | n.d        |
| Er              | 11.1  | 12.1  | 11.2  | 17.6  | n.d        |
| Tm              | 0.38  | 0.80  | 0.91  | 0.55  | n.d        |
| Yb              | 9.4   | 9.5   | 8.9   | 14.8  | 2.4        |
| Lu              | 1.36  | 1.38  | 1.30  | 2.1   | 0.4        |
| Hf              | 23    | 32    | 15.5  | 66    | 4.3        |
| Ta              | 10.4  | 18.0  | 7.7   | 36    | 0.9        |
| W               | 4.4   | 6.7   | 1.3   | 2.8   | <1         |
| Pb              | 37    | 15.0  | 15.5  | 67    | 6.5        |
| Th              | 30    | 45    | 28    | 89    | 8.8        |
| Hg (<|dl)      |       |       | <|dl|  <|dl|  0.0      |
| U               | 8.0   | 7.6   | 1.98  | 9.2   | 2.1        |

LOI = Loss of ignition (wt.%); <|dl| = < to detection limits; n.d = not determined;
5. Discussions

5.1. Genesis of the Studied Geophagic Clayey Materials

Sabga geophagic clayey materials outcrop within trachyte. Determining the origin of geophagic clayey materials (meteoric weathering or hydrothermal alteration) for pharmaceutical use is essential because the suitability of a clay deposit as pharmaceutical excipient greatly depends on its geological nature (sedimentary, residual, and hydrothermal) and mineral composition of the deposits, which have an important effect on texture and particle size distribution, and consequently, on the rheological properties (flow) of the powder mass [59, 31]. It is relatively easy to establish an hydrothermal origin of high temperature facies, with potassic alteration for example [37]. As the temperature of transformation decreases, there is a great resemblance between meteoric and hydrothermal minerals. The assemblages can have the following phases in common: smectite, vermiculite, kaolinite and illite. However, the geological setting, field characteristics of the deposit, associated minerals and geochemical signatures allow to distinguish differences between the two alteration phenomena [54, 20, 21, 65]. To determine the origin of Sabga geophagic clayey materials, these geological features are taken into account. In Cameroon, where several localities abound in edible clayey material deposits developed during meteoric alteration of various rocks. Among the different clay minerals, we have friable halloysite deposit resulting from weathering of trachyte and phonolite from Mount Bamenda at Santa, Bali, Bambili and Sabga areas [69], more than 15 m thick kaolinitic clays at Lembo and Bana, which may have developed during meteoric alteration of various rocks (granite, rhyolite, trachyte and basalt) of the Bana tertiary ring complex [94, 65] and kaolin deposit of Balengou (halloysite) which may have been formed from alteration of trachyte [63].

The geological setting of Sabga area is a volcanic environment, composed of trachyte, rhyolite, basalt and ignimbrite (Figure 2). These rocks have different characteristics as rocks at high temperature and pressure like onycolites. This zoning morphology is characterised by the absence of tectonic structures such as veins, foliation and fault. From the above details, it is suggested that the process of forming geophagic clayey materials is completely different from the typical hydrothermal alteration profile [10, 11, 65].

With these geological features, the field characteristics are close to those generally reported for meteoric weathering clayey materials. According to various analytical techniques (XRD, FTIR, scanning electron microscopy and chemical analyses), Sabga geophagic clayey materials have the following mineral phases: smectite + kaolinite, + feldspar + quartz ± goethite ± hematite. This paragenesis also indicates that the smectite and kaolinite are the alteration product of feldspar.

In humid tropical climate, meteoric weathering generally leads to the formation of kaolinite with poor crystallinity due to numerous impurities as Fe, Ti, Cr and Ni in kaolinite structure [71, 10, 11, 65]. No hydrothermal alteration mineral indicator as florencite, pyrite, chrysotile and lizardite or any ore associated minerals was found in the geophagic clayey materials [10, 11, 65, 66]. Sabga geophagic clayey materials contain some amounts of alkalis and alkali earths despite their location on steep slope hills under tropical humid climate. These are good drainage conditions that must have favoured the leaching of these elements during meteoric weathering [80].

Geochemical compositions can give information about the origin of the alteration. From the oxides chemistry, the small TiO$_2$ contents and the high Fe$_2$O$_3$ contents corroborate with meteoric weathering because meteoric weathering generally enhances the presence of ferric oxides and hydroxides in tropical climate [54, 74, 65]; and TiO$_2$ may be concentrated in anatase by hydrothermal fluids [58, 34]. Ce anomalies are well known in weathering and lateritic soils formation process.

The studied geophagic clayey materials were compared to the REE composition of Sabga B08 trachyte of [42] and of chondrite CI [52]. In Sabga geophagic clayey materials, negative Ce and Eu anomalies have been observed in all patterns (Figure 9a, b). The REE patterns are parallel, suggesting that all the geophagic clayey materials were formed through the same process. Generally, Positive Ce anomalies are well known in weathering and lateritic soils formation process. These anomalies are produced when Ce$^{3+}$ is oxidized to Ce$^{4+}$ favoured under an oxidizing environment in the weathering profile [55, 10, 11, 19, 15, 67, 65]. Conversely, hydrothermal processes usually occur in more reducing conditions [20, 21] where Eu$^{3+}$ is reduced to Eu$^{2+}$ and fractionates differently from other REEs. Commonly, the absence of positive and negative Ce anomaly confirms hydrothermal alteration [65, 79]. Figure 9 points out Ce anomalies, as it is the case of supergene smectite and halloysite clay deposit of Lisbonne in Portugal [73], clay minerals of Kombelcha and Bombowha in Ethiopia [28], quartz-rich smectitic clays of the northern part of Cameroon formed in a confined environment under a Sahelian climate [62] and halloysite clay deposit of Balengou [80].

Geophagic clayey materials samples from Sabga display peculiar deep negative Ce and Eu anomalies (Figure 9). These anomalies reflect a selective leaching of these two elements during geophagic clayey materials formation processes. The chondrite-normalized REE patterns of the geophagic clayey materials show LREE enrichments and a negative Eu anomaly (Figure 9b). The HREE patterns are almost flat (Figure 9b). Eu anomaly suggesting that it developed at least partially in response to feldspar weathering, where most of the Eu is hosted [7]. These characteristics are similar to that of clay-rich sediments from central Uganda Makoro derived from granitoid rocks [68].
5.2. Medical Interest of the Studied Geophagic Clayey Materials

The geochemical and mineralogical constitutions of these soils are likely to have some implication on the health of the geophagic individuals. Table 3 indicates that the chemical composition range of geophagic clayey materials from Sabga is very similar to that of natural smectite used in pharmacy and cosmetics [46] except for its high Fe₂O₃ contents. Sabga geophagic clayey materials have radioactive, carcinogenic and teratogenic (or birth defects) elements, higher than those reported in recommended pharmaceutical clays [51]. Thus from the chemical compositions, the direct ingestion of Sabga geophagic clayey materials may be hazardous. The raw material should be purified before pharmaceutical and cosmetic uses.

Samples SA, SD with relatively higher amounts of smectite, sample SB, SC with a combination of kaolinite and smectite could contribute to the alleviation of gastrointestinal discomfort related to diarrhea and the elimination of toxins in geophagic individuals. According to [30] oxides of Fe would result in different pigment colours in soils including reddish and yellowish colours (Figure 3). Most of the studied samples were reddish and yellowish and hematite and goethite were identified in some of them. This may be attributed to the presence of Fe-bearing mineral (Table 1) and Fe oxides and hydroxides in the soils (Table 3). Geophagic individuals from Sabga locality indicated that they ingested the soils for famine and pregnancy related reasons. Despite Fe providing supplement for those who lack iron in their blood, a high level of Fe in the blood could result in hemochromatosis and possible death [91]. Besides, the consumption of Fe-rich lateritic soils may negatively influence the utilisation of copper and zinc in the body [9].

Zinc concentration in the studied Sabga geophagic clayey materials were highest in samples SA, SB, SC, SD with respectively 248, 147, 177 and 989 ppm. Having a recommended value for zinc of 0.01ppm [91], excessive consumption of these clayey materials may lead to unusual drowsiness, nausea, vomiting, diarrhea and growth retardation of an unborn baby [75]. Zinc plays a vital role in cellular metabolism and is a major component of body tissues and fluids [78]. Most catalytic activities of enzymes require zinc, which is equally required for immune function, wounds healing, protein synthesis, DNA synthesis and cell division. Zinc ions are effective antimicrobial agents even at low concentrations [70]. Zinc is also required for proper functioning of the senses of taste and smell and supports
normal growth and development during pregnancy, childhood, and adolescence [4]. There are claims of zinc having antioxidant properties, which may protect against accelerated aging and helps speed up the healing process after an injury. However, studies differ as to its effectiveness [70].

Chromium has been reported to be responsible for lung cancer and death [76]. The highest concentration of Cr was recorded in all samples as shown in Table 3 (1.9-7.0 ppm) which is above the World Health Organization recommended value of 0.05 ppm [91]. Sample SD had the highest concentration of Pb with 67 ppm, SB and SC have the lowest concentration of 15.0 and 15.5 ppm respectively. Daily recommended value for lead (Pb) is 0.01 ppm [91]. The presence of Pb in the samples poses a problem as there is no prescribed limit for its consumption due to its extreme toxic nature. Previous studies revealed that exposure to lead may cause intelligence decline in children and as well as cancer in adults [90]. Consumption of these clayey materials over a long period may result in serious health effects such as coma, seizure, dysfunctioning of the kidney, liver and heart of the consumers.

The particle size diameter of geophagic clayey materials (Table 2) was lower than 10 μm. The PSD curves for the samples are very close to each other (Figure 8). Knowledge of particle size distribution features of the raw materials is essential for the right pre-formulation steps of cosmetic and pharmaceutical products [39]. Based on the literature [72] the powder particle size distribution applicability can vary. Finer powders have higher skin adhesion and provide better softness when applied on skin. One example showing that particle size influences powder properties is the research published by [72], which demonstrated that products for topical application containing in their composition clay of particle size in the range of 2 and 9 μm. This reduced particle size range suggests the application of clays in cosmetics. According to literature, particles smaller than 63 μm may have anti-inflammatory effects and may assist in the skin hydration, retaining moisture due to the high skin adhesiveness [16]. Samples from Sabga had higher mean values of D (90) 4 - 60 μm (Table 2) compared with 8.3 – 53.8 μm for samples of pharmaceutical clays originated from the sand extraction residue of mining activities in the hinterland of São Paulo state, Brazil [39].

6. Conclusion

Prospection and characterization studies carried out in the Sabga locality have revealed the presence of heterogeneous geophagic clayey materials along the higher slopes of Sabga hill formed by trachyte. It comprises smectite and kaolinite resulting from the alteration of trachyte. Non-clay mineral assemblages comprise feldspar, quartz, goethite and hematite, pointing out an alteration controlled by oxidizing environment. No hydrothermal alteration mineral indicators are identified in geophagic clayey materials. Trachyte and chondrite normalized trace element patterns for representative geophagic clayey materials show negative Ce and Eu anomalies. All these characteristics lead to the interpretation that geophagic clayey materials are from a meteoric weathering origin. From granulometric, mineralogical and chemical properties, geophagic clayey materials can be considered as pharmaceutical clays after some previous treatment to reduce the amount of toxic metals and crystalline silica (quartz). Studies on bioavailability of Fe and Zn elements in the gastrointestinal tract are however necessary to determine which percentage of the total concentration is actually available for assimilation upon ingestion. Nevertheless, the potential medicinal application of these clayey materials is supported by the smectite and kaolinite contents, and fines particles size. In contrast, the trace elements are pointers of probable adverse effects on humans.

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