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Comprehensive physicochemical study of dioctahedral palygorskite-rich clay from Marrakech High Atlas (Morocco)

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Abstract This study is devoted to the physicochemical and mineralogical characterizations of palygorskite from Marrakech High Atlas, Morocco. The raw clay and its Na⁺-saturated <2 μm fraction were characterized using chemical, structural, and thermal analytical techniques. Measurements of specific surface area and porous volume are reported. The clay fraction was found to be made up of 95 % of palygorskite and 5 % of sepiolite. An original feature of this palygorskite is its deficiency in zeolitic H₂O. The half-cell structural formula of its dehydrated form was determined on the basis of 21 oxygens to be (Si_{7.92}Al_{0.08})(Mg_{2.15}Al_{1.4}Fe_{0.4}Ti_{0.05}□₁)(Ca_{0.03}Na_{0.08}K_{0.04})O₂₁, while the hydrated form could be formulated as (Si_{7.97}Al_{0.03})(Mg_{2.17}Al_{1.46}Fe_{0.40}Ti_{0.05})(Ca_{0.03}Na_{0.07}K_{0.03})O_{20.18}(OH)_{1.94}(OH₂)_{3.88}·2.43 H₂O. These formulas show that the (Al³⁺+Fe³⁺)/Mg²⁺ ratio is around 0.84, revealing a

pronounced dioctahedral character. Further, inside its octahedral sheet, it was determined that the inner M₁ sites are occupied by vacancies, whereas the M₂ sites are shared between 90 % of trivalent cations (78 % for Al³⁺ and 22 % for Fe³⁺), 7.5 % of Mg²⁺, and 2.5 % of Ti⁴⁺, all of them linked to 1.94 of structural hydroxyls. The two remaining Mg²⁺ by half-cell occupy edge M₃ sites and are coordinated to 3.88 molecules of OH₂. Channels of this palygorskite are deficient in zeolitic H₂O since they contain only 2.43 H₂O molecules. A correlation was found between these results and the observation of very intense and well-resolved FTIR bands arising from dioctahedral domains (mainly Al₂OH, Fe₂OH, and AlFeOH) along with very small responses from a trioctahedral domain (Mg₃OH). Accordingly, a schematic representation of the composition of the octahedral sheet was proposed. The cation exchange capacity, specific surface area, and total pore volume were also assessed to be ca. 21.2 meq/100 g, 116 m²/g, and 0.458 cm³/g, respectively.

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Introduction

Palygorskite, also called attapulgite, is an interesting fibrous hydrated aluminum magnesium silicate mineral with versatile industrial applications (Murray 2000). From historical and cultural viewpoints, palygorskite was known by ancient civilizations since it was used during Pre-Columbian Mesoamerican period to manufacture the prominent and mysterious hybrid pigment, called Maya Blue, which is a combination of palygorskite, indigo dye, and metallic particles (Van Olphen 1966; Yacamán et al. 1996;

Polette-Niewold et al. 2007). Most recently, increasing interest was devoted to fibrous clay minerals due to their adsorptive capacity and their ability to be combined to organic or inorganic compounds, leading to the design of versatile nanostructured functional materials (Zhao et al. 2006; Bouna et al. 2011; Su et al. 2012; Liu et al. 2012; Li et al. 2012). In comparison with other clay minerals, palygorskite is a fuller's earth exhibiting specific desirable sorptive, colloidal-rheological, and catalytic properties (Galán et al. 1994; Galán 1996; Wang et al. 2012). Technological applications of palygorskite were shown by Galán (1996) to be due to its physicochemical properties, mainly composition, surface area, porosity, among others, and especially fibrous structure.

In this context, the structure of palygorskite was described by Bradley's model (Bradley 1940; Bailey 1980). It consists of continuous sheets of tetrahedrons (T) in which the apical oxygens are alternatively inverted pointing in opposite directions and binding to a discontinuous octahedral sheet (O), which forms ribbons elongated in the direction of the *c*-axis (Krekeler and Guggenheim 2008; Fig. 1a, b). In this clay mineral, the TOT ribbons have an average width along the *b*-axis of two linked tetrahedral chains. Besides, they are disposed according to a chessboard-like arrangement when observed on the (001) plane, linked one another via Si–O–Si bridges. The octahedral ribbon is made up of five sites whose the inner one is M_1 , the two following ones (M_2) are coordinated to two hydroxyls, and the last two located on edges are M_3 (Fig. 1c). The linked ribbons thus form a continuous 2:1 layer along the *c*-axis (Fig. 1a, c) with a limited extent along the *b*-axis (Fig. 1b). Rectangular channels of about $3.7 \times 6.0 \text{ \AA}^2$ wide (Huang et al. 2007) are present. They lie between the blocks of opposing 2:1 ribbons and extend parallel to the fiber direction. As a result of the discontinuity of the silica sheets, silanol groups (Si–OH) are present at the external surfaces of the particles located at the edges of the channels (Fig. 1) (Shariatmadari 1998; Ruiz-Hitzky 2001). Further, the clay mineral contains two kinds of water molecules, ones coordinated to the octahedral cations and the other weakly bonded in the channels, namely the zeolitic H_2O . These channels may also contain exchangeable cations.

Isomorphous substitutions principally of Al and Fe for Mg occur in the octahedral sites of this silicate. Thus, a small number of exchangeable compensating Na^+ , K^+ , Ca^{2+} , and Mg^{2+} cations (García-Romero et al. 2007), zeolitic H_2O , and four structural OH_2 molecules per formula unit bonded to M_3 edge sites are contained in the channels (Mifsud et al. 1978; Suárez and García-Romero 2006). Palygorskite has a general theoretical formula $(Mg_{5-y-z}R^{3+}_y\Box_z)(Si_{8-x}R_x^{3+})O_{20}(OH)_2(OH_2)_4R_{(x-y+2z)/2}^{2+}(H_2O)_4$, where \Box is a vacant site and R represents Al or Fe, depending on the origin and deposition conditions (Huang et al. 2007). So, a variable dioctahedral-trioctahedral

character can be assigned to palygorskite (Galán and Carretero 1999; Chahi et al. 2002). The dioctahedral end-member at $y = 2$ and $z = 1$ is characterized by the formula $(Mg_2R^{3+}_2\Box_1)(Si_{8-x}R_x^{3+})O_{20}(OH)_2(OH_2)_4R_{x/2}^{2+}(H_2O)_4$ showing that only four of five octahedral positions available are filled with R^{3+} occupying the M_2 sites and Mg^{2+} occupying the M_3 while M_1 is vacant (Güven et al. 1992). On the other hand, the trioctahedral end-member palygorskite at $y = 0$ and $z = 0$ can be represented by the formula $(Mg_5)(Si_{8-x}R_x^{3+})O_{20}(OH)_2(OH_2)_4R_{x/2}^{2+}(H_2O)_4$ for which all the five octahedral positions are filled with Mg^{2+} .

The occurrence of palygorskite in Marrakech High Atlas was reported by several authors (Daoudi and Deconninck 1994; Ben Aboud et al. 1996; Daoudi 2004). These studies were mainly focused on the knowledge of their genesis and/or their particular geological features. Likewise, we recently demonstrated the possibility of immobilization of TiO_2 nanoparticles ($\sim 8 \text{ nm}$) on fibers of this palygorskite to produce nanocomposite materials as original supported photocatalysts. Indeed, they exhibited remarkable photocatalytic activity in the removal from aqueous medium of Orange G dye widely used in textile industry (Bouna et al. 2011).

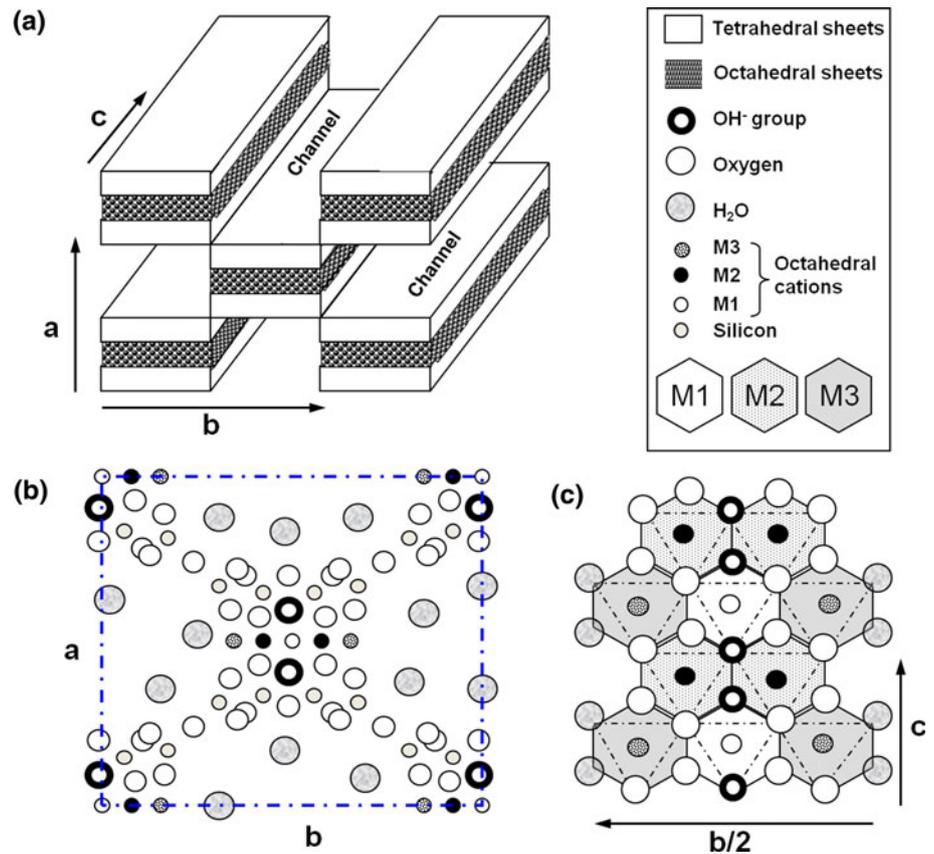
Despite all these studies, a thorough characterization of palygorskite from Marrakech High Atlas is still incomplete. Pursuing a general program of research on the characterization, properties, and technological development of natural clays from Morocco (Rhouta et al. 2008; Bouna et al. 2011, 2012; Ait Aghzzaf et al. 2012), this study aims at a detailed analysis of the raw clay from Marrakech High Atlas and its $<2 \mu\text{m}$ fraction in order to determine their mineralogical, physicochemical, textural, and structural properties, especially by elucidating the composition of its octahedral sheets. Knowledge of these properties is vital for further considering the clay for use as a cheap abundant natural starting material in the development of innovative clay-based materials.

Geological setting

The Uppermost Cretaceous-Paleogene sediments of the Marrakech High Atlas consist chiefly of alternations of marl, carbonate, gypsum, and phosphatic bed suggesting a shallow sea environmental deposit (Marzoqi 1990). Sedimentary facies and clay assemblages' evolution of this series is studied in the Asni region, 45 km South of Marrakech, where they constitute thick accumulations exceeding 400 m. The sedimentary sequences are organized in two systems which are separated by a regional-scale discontinuity (Fig. 2):

- (i) The lower sedimentary system which is probably of Maastrichtian age (Chellai et al. 1995), comprise elementary sequences composed by alternations of

Fig. 1 Schematic representations of palygorskite structure: **a** overview of the fibrous morphology, the presence of channels, and the formation of discontinuous ribbons; **b** projection on (001) plane; and **c** projection on (100) plane showing an octahedral ribbon with the M₁, M₂, and M₃ sites as well as structural hydroxyls, coordinated and zeolitic H₂O [Suárez and García-Romero (2006)]



gypsum, occurring as bands and veins, and banks of bioclastic carbonates which became increasingly diversified upwards. The clay fraction is mainly composed of smectite, illite, and little amount of palygorskite (from Pk0 to Pk1 levels).

- (ii) The upper sedimentary system is of Paleocene to Lutetian age (Chellai et al. 1995). The lower member is composed of yellow heterogeneous marls followed by dolomitic bars and phosphate deposits. The associated clay fraction is composed essentially by fibrous clay (palygorskite and sepiolite) quite in the same proportions (Pk2, Pk3). The upper member consists of a thick series of marls intercalated by thin dolomitic or calcarenite bars. This unit is characterized by the widespread occurrence of palygorskite, which reaches its maximum at the top of the series (Pk7). Thus, this mineralogical and physicochemical study is focused on this PK7 palygorskite-rich unit.

treated with HCl solution (0.2 N). The almost pure Na⁺-homoionic clay fraction (labeled Na⁺-PK7) was recovered by the procedure described elsewhere (Rhouta et al. 2008). For removing poorly crystalline iron-rich impurities, which could be present as iron oxides and/or iron hydroxide gel (Goodman et al. 1976), the Na⁺-exchanged <2 μm fraction (Na⁺-TAG) was treated with a dithionide-citrate solution buffered with sodium bicarbonate according to the procedure reported by Mehra and Jackson (1956). For this purpose, 1 g of <2 μm fraction powder was dispersed under stirring at 80 °C in an aqueous solution of 40 mL of sodium citrate (0.3 mol/L) and 5 mL of sodium bicarbonate (1 mol/L). Thereafter, 500 mg of sodium dithionide was added to this dispersion which was further kept at 80 °C under stirring for additional 15 min. Finally, the clay material was recovered by centrifugation at ×604 for 10 min and washed several times to remove excess of reacting compounds.

Experimental

Purification

Raw clay samples (designated PK7 where PK stands for “Plateau Kik”) were first crushed and sieved to 50 μm and

Characterizations

Microstructural observations and qualitative compositions were performed on raw clay and its Na⁺-exchanged <2 μm fraction with a Jeol JSM 6400 scanning electron microscope (SEM) and a Jeol JEM 2010 transmission electron microscope (TEM) equipped with an energy-dispersive

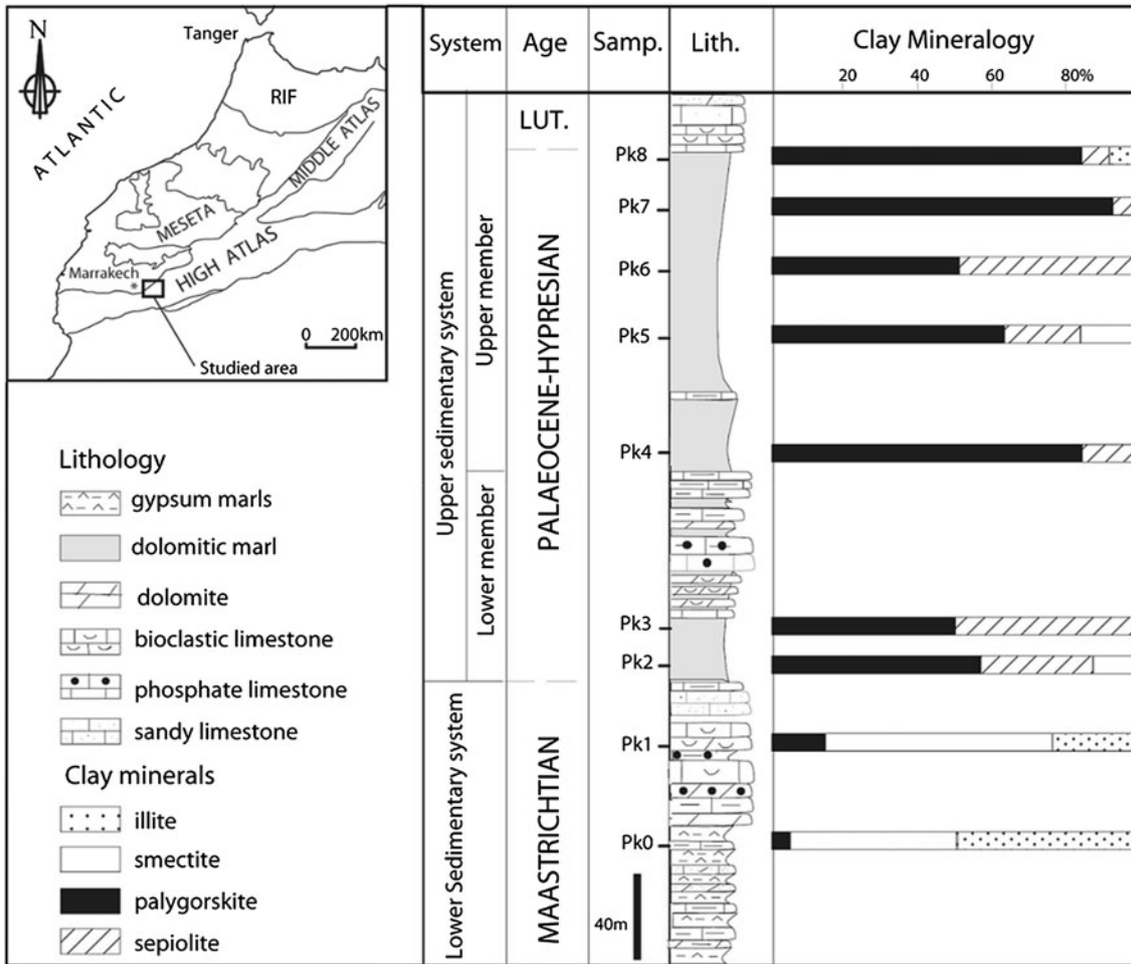


Fig. 2 Geographical location of the sampling area (top left corner). Lithology and clay mineralogy of the Asni section

X-ray Tracor analyzer (EDS). SEM observations were performed by spreading and fixing the powder sample on double-sided adhesive and conductive tape stuck on the sample holder. To ensure a good electron conduction, the prepared samples were coated by cathodic sputtering with a thin silver film (a few nanometers of thickness) using an Edwards S150B metalizer prior to be introduced in the microscope. For TEM examinations, a small amount of material powder was first dispersed in ethanol with an ultrasonic bath. After that, droplets of this dispersion were placed on a copper grid (3 mm in diameter) covered by a thin layer of amorphous carbon. After evaporation of ethanol, particles of the sample were retained on the grid and it was introduced in the TEM apparatus using a sample holder.

X-ray diffraction (XRD) patterns were recorded for raw clay (PK7) and $<2 \mu\text{m}$ Na^+ -saturated fraction (Na^+ -PK7) on oriented flat plates over the 2θ angular range $2\text{--}60^\circ$. A Seifert XRD 3000TT apparatus equipped with a diffracted beam graphite monochromator was used in the Bragg–Brentano configuration ($\text{Cu K}_{\alpha(1+2)}$ radiation). The

thermogravimetric (TG) and differential thermal analysis (DTA) thermograms were collected using a Setaram Lab-sys apparatus using sample masses of ca. 60 mg and a temperature ramp of 10 K/min from ambient to 1,273 K. Solid-state ^{27}Al and ^{29}Si magic angle spinning nuclear magnetic resonance (MAS-NMR) spectra were collected at room temperature with a Bruker Advance 500 spectrometer equipped with a 4-mm cross-polarization magic angle spinning probe fitted with a standard spinning assembly. ^{27}Al MAS-NMR spectra were obtained at 130.32 MHz, while ^{29}Si MAS-NMR spectra were recorded at 99.36 MHz. ^{27}Al chemical shifts were recorded with respect to $[\text{Al}(\text{H}_2\text{O})_6]^{3+}$ as an external reference, and ^{29}Si chemical shifts were referred to tetramethylsilane (TMS).

The chemical compositions of crude clay and its $<2 \mu\text{m}$ Na^+ -saturated fraction were determined by inductively coupled plasma (ICP) by mean of a Thermo Electron Series X2 spectrometer equipped with a Meinhard nebulizer and a SimulScan FTE-type detector. The clay samples were heated to 800°C overnight prior to analysis to dehydrate the samples (Caillere et al. 1982; Rautureau et al. 2004).

Mid-infrared (MIR) spectroscopic studies were undertaken using a Nicolet 5700 spectrometer under ambient conditions in the frequency range $400\text{--}4,000\text{ cm}^{-1}$ using KBr pellets. The pellets contained a mixture of 2 mg of clay material with 198 mg of KBr ($\sim 1\text{ wt } \%$). The near-infrared (NIR) spectra were obtained using a Perkin Elmer Lambda 19 spectrophotometer equipped with an integrating sphere coated by BaSO_4 , which was used as a reference. The spectra were recorded with samples deposited on quartz plates from $4,000\text{ to }8,000\text{ cm}^{-1}$ at a resolution of 5 cm^{-1} .

The cation exchange capacity (CEC) of the raw clay (PK7) and its purified Na^+ -exchanged fraction (Na^+ -PK7) was determined according to the method based on adsorption of hexamine cobalt complex (Mantin and Glaeser 1960). To improve clay minerals particles to be sufficiently wetted by the adsorbate, material dispersion was stirred under ultrasounds to disaggregate clay mineral fibers. The specific surface area was measured from the adsorption–desorption isotherms of N_2 at 77 K using an ASAP 2020 V 3.01 H adsorption analyzer from Micromeritics. Prior to N_2 adsorption–desorption measurements, samples weighting 0.162 g were outgassed either at $150\text{ }^\circ\text{C}$ overnight or for 24 h or at $140\text{ }^\circ\text{C}$ for 9 h till the pressure reached around 10^{-2} Torr. The specific surface area was determined by the Brunauer–Emmett–Teller (BET) method (Brunauer et al. 1938).

Results and discussion

Microstructural analysis (TEM–SEM–EDS)

TEM micrograph obtained from raw clay shows microfibrils mixed with rhomboidal crystals characteristic of carbonates (Fig. 3a). Punctual EDS analysis performed on selected area of several fibers or along a same fiber (spectra not reported) revealed that these fibers are aluminum magnesium silicate with noticeable amount of iron.

The overview of Na^+ -exchanged fine fraction afforded by SEM imaging (Fig. 3b) shows fibers bundles free from carbonate impurities. The fibers appear either as well-separated individual fibers or as assemblages of several fibers having an average length of ca. $1\text{ }\mu\text{m}$ and a diameter ranging from 10 to 50 nm. Besides, these micrographs exhibit various sizes of interfibrils pores, widely ranging from 10 nm to $1\text{ }\mu\text{m}$, indicating the occurrence of especially mesopores ($2\text{ nm} < \text{size} < 50\text{ nm}$) and macropores ($\text{size} > 50\text{ nm}$) (Rouquerol et al. 2003). But, this does not exclude the presence of interfibrils micropores ($\text{size} < 2\text{ nm}$) (Rouquerol et al. 2003, Cases et al. 1991) which could not be identified due to magnitude limitation of the TEM apparatus used herein.

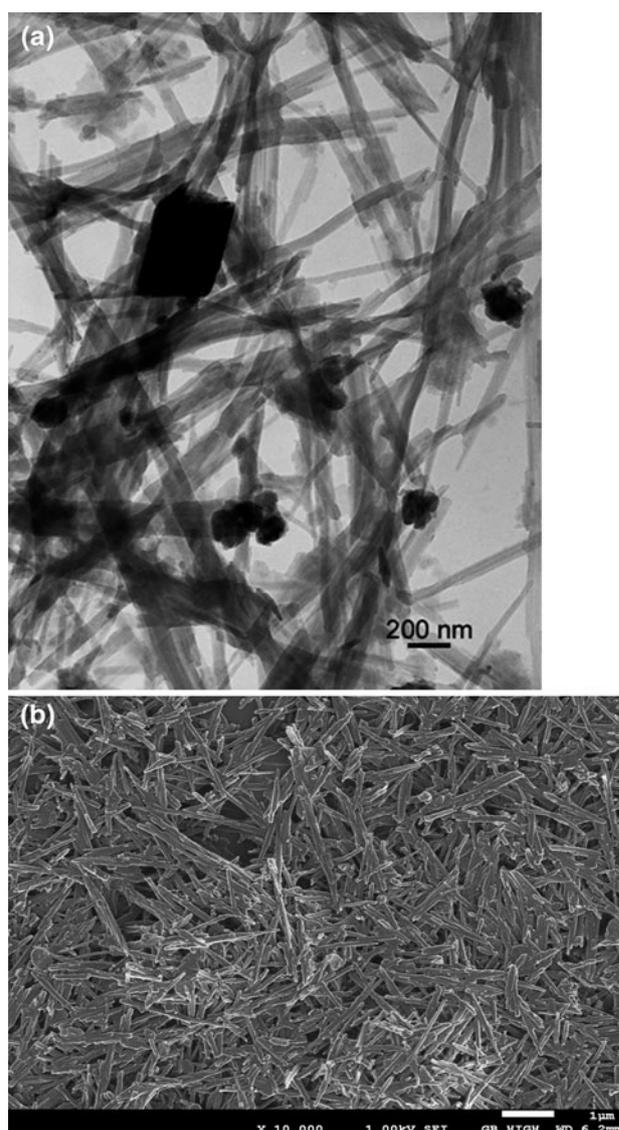


Fig. 3 **a** TEM micrograph of raw clay (PK7) showing clay mineral fibers mixed with rhomboidal crystals of carbonates. **b** SEM micrograph of Na^+ exchanged purified fine fraction (Na^+ -PK7)

Crystallographic structure and phase identification (XRD)

The XRD patterns of raw clay sample PK7 (Fig. 4a) show the reflections of a fibrous clay mineral (P) along with those corresponding to quartz (JCPDS file: 03-065-0466) and to two kinds of carbonates: calcite (C) $\text{Mg}_{0.03}\text{Ca}_{0.97}\text{CO}_3$ (JCPDS file: 01-089-1304) and ankerite (A) $\text{Ca}_{1.01}\text{Mg}_{0.45}\text{Fe}_{0.54}(\text{CO}_3)_2$ (JCPDS file: 01-084-2066) in good agreement with TEM–EDS analyses. Upon purification, carbonates peaks disappear, whereas those of quartz are still present indicating the success of the removal of carbonate while minor amount of quartz remains in the sample (estimated to only a few percent from the relative XRD intensity). At the

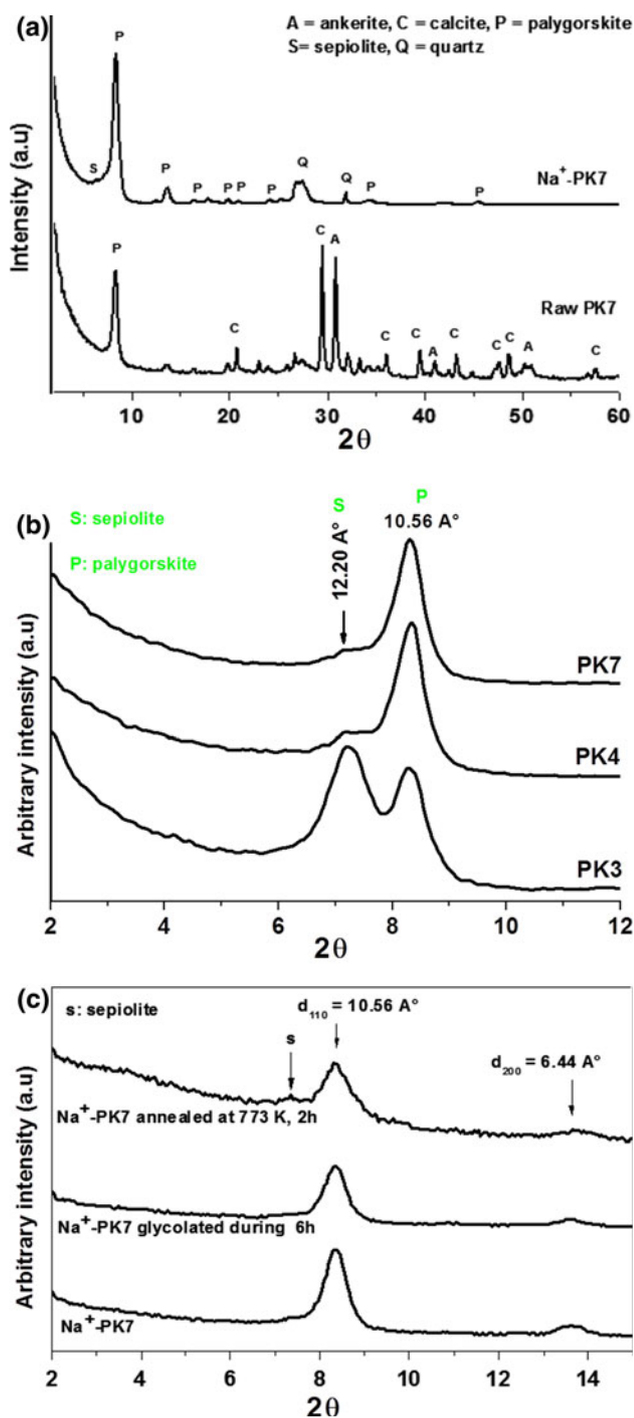


Fig. 4 XRD patterns of the raw clay (PK7) and its isolated fine fraction (Na^+ -PK7): **a** before (PK7) and after purification (Na^+ -PK7); **b** Comparison of XRD patterns of samples picked up from different units (PK3, PK4, and PK7); **c** XRD patterns in the small angle region of fine clay fraction (Na^+ -PK7) before and after successive different treatments. Reproduced with permission from Bouna et al. (2011)

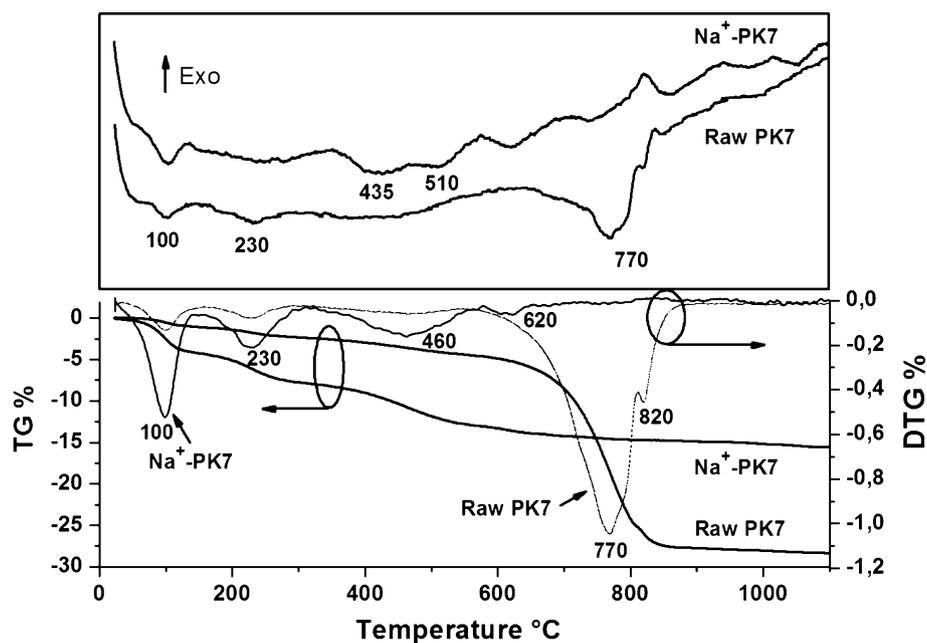
same time, reflections of fibrous clay mineral appear more resolved and intensified. Their angular positions are quite similar to those reported for palygorskite from other

occurrences (Christ et al. 1969; Chisholm 1990, 1992; Artioli and Galli 1994; Fernández et al. 1999). The d_{110} reflection found around 10.56 \AA ($2\theta \approx 8.36^\circ$) confirms that it corresponds to palygorskite. Sepiolite coexists with palygorskite in the PK7 clay but in a minor amount as evidenced by the detection of a small shoulder at around 12.2 \AA ($2\theta \approx 7.25^\circ$) corresponding to its (110) peak (Fig. 4a). The intensity of this additional peak is found to increase for the samples taken from deeper units (PK3 and PK4) as evidenced by XRD (Fig. 4b). Semiquantitative estimations, based on “peak height ratios” (Thorez 1976; Holtzapffel 1985; Pevear and Mumpton 1989; Galhano et al. 1999) of d_{110} reflections of palygorskite and sepiolite, deduced from X-ray diagram of Na^+ -PK7 sample (Fig. 4a, b), indicate that the fibrous clay mineral fraction is a mixture of 95 % of palygorskite and 5 % of sepiolite in good agreement with geological context described above. The diffractograms obtained from oriented flat plates of the purified Na^+ -exchanged fine (<2 μm) fraction (Na^+ -PK7), after being either glycolated for 6 h or heat treated at $500 \text{ }^\circ\text{C}$ for 2 h (Fig. 4c), show that the d_{110} reflection at 10.56 \AA is not affected by ethylene glycol solvation or the heat treatment. This confirms that the palygorskite investigated herein is not associated to any swellable smectite in contrast to palygorskite from phosphorite deposits of Gantour, Morocco (Chahi et al. 2002) or the palygorskite from other occurrences (Gionis et al. 2006, 2007).

Thermal behavior (TG-DTA)

Evidence of the removal of carbonates is given by the disappearance in TG-DTA thermograms of purified fraction of the complex endothermic event observed for raw clay in the temperature range $560\text{--}885 \text{ }^\circ\text{C}$ associated with a weight loss of about 23 %. TG-DTA thermograms of Na^+ -exchanged fraction (Fig. 5) are typical to those reported in previous studies (Giustetto and Compagnoni 2011). They show an endothermic peak at about $100 \text{ }^\circ\text{C}$ associated with a mass loss of 4.4 wt% generally ascribed to the removal of both physisorbed water and a part of zeolitic H_2O . This weight loss is lower than the 6 wt% recorded for palygorskite from Montestrutto, Italy (Giustetto and Compagnoni 2011) and the 9 wt% reported for other palygorskite (Mifsud et al. 1978). This is probably due to the removal of a low amount of physisorbed water upon samples drying at $80 \text{ }^\circ\text{C}$ prior to thermal analysis. This might also indicate that this weight loss predominantly corresponds to zeolitic H_2O and confirms a deficiency in zeolitic H_2O of the palygorskite from Marrakech High Atlas. Another broad endothermic effect, observed between 160 and $300 \text{ }^\circ\text{C}$ with a maximum at around $230 \text{ }^\circ\text{C}$, is accompanied with a mass loss of about 3.6 wt%. It is due to the removal of remaining zeolitic H_2O with the first fraction of Mg-coordinated OH_2 (M_3 site). This temperature is higher than that ($\approx 205 \text{ }^\circ\text{C}$)

Fig. 5 Thermal gravimetry (a) and DTA thermograms (b) of raw clay (PK7) and its fine fraction (Na^+ -PK7)



reported by Gionis et al. (2006) for palygorskites Gr1 from Pefkaki in Greece and PFI-1 originating from Gadsden County, Florida (USA). This difference could be explained by the presence of ca. 5 % of sepiolite associated to palygorskite investigated herein, as revealed by XRD analysis, because this endothermic effect is found at higher temperature (>250 °C) for sepiolite (Frost and Ding 2003; Giustetto et al. 2011). The global weight loss (≈ 8 %) recorded in the low-temperature range ($T < 300$ °C) is lower than previously reported values, for example, 12.5 % (Giustetto and Chiari 2004) and 13.7 % (Artioli et al. 1994), implying the occurrence of H_2O deficit, mainly zeolitic H_2O , in our samples (Mifsud et al. 1978, Giustetto and Compagnoni 2011). The zeolitic H_2O deficiency seems to be more pronounced in Marrakech High Atlas palygorskite by comparison with the palygorskite from Montestrutto, Italy (Giustetto and Compagnoni 2011: ≈ 9 wt%).

Between 330 and 590 °C, a broad endothermic peak, centered around 460 °C, associated with mass loss of about 5.2 % is attributed to the removal of the residual fraction of Mg-coordinated OH_2 (Giustetto and Chiari 2004; Giustetto and Compagnoni 2011), which causes the collapse and the formation of “palygorskite anhydride” (Preisinger 1963 in Giustetto and Compagnoni 2011). In the high-temperature region ($T > 600$ °C), an endothermic effect peaking at 620 °C associated with supplementary weight loss around 2 % is due to dehydroxylation resulting in the formation of clinoenstatite (Hayashi et al. 1969 in Giustetto and Compagnoni 2011). Finally, an exothermic effect observed at about 820 °C is due to the crystallization of a new anhydrous mineral phase.

According to Caillère and Hénin 1961, the weight losses described above could be assigned to different kinds of water of palygorskite according to temperature ranges by assuming that zeolitic H_2O is lost below 200 °C, structural OH_2 is lost between 250 and 400 °C, and hydroxyl groups above 400 °C. Nevertheless, other authors stated that the partitioning of these mass losses among different kinds of water is not so strict (Giustetto and Chiari 2004; Giustetto and Compagnoni 2011). As suggested by Giustetto and Compagnoni (2011), the 2 wt% decrease at high temperatures can indisputably be ascribed to the loss of framework hydroxyls, and 8 wt% among the remaining weight loss (≈ 13 wt%) can be reasonably attributed to structural OH_2 , while 5 wt% may be related to zeolitic H_2O .

Tetrahedral versus octahedral occupation of Si^{4+} and Al^{3+} (MAS-NMR)

MAS-NMR spectroscopy did not reveal differences between the raw clay and its fine fraction. Typical ^{27}Al -NMR spectrum (Fig. 6a) shows two peaks at 4.1 and 71.4 ppm corresponding to octahedral and tetrahedral Al, respectively (Woessner 1989; Komarneni et al. 1986a, b). The peak at 4.1 ppm appears significantly more intense than the peak at 71.4 ppm, which indicates, as expected, that Al principally occupies octahedral sheet, whereas only 5 % of Al^{3+} , deduced from peaks intensities ratio, occupies tetrahedral sheet. Typical ^{29}Si -NMR spectrum (Fig. 6b) reveals a doublet of peaks at -91.7 and -96.8 ppm corresponding to two nonequivalent crystallographic sites of tetrahedral Si cations Q3(0Al) (Komarneni et al. 1986a, b). A small peak at -84.9 ppm likely corresponds to component Q3(1Al) (Komarneni et al. 1986a) confirming the substitution in

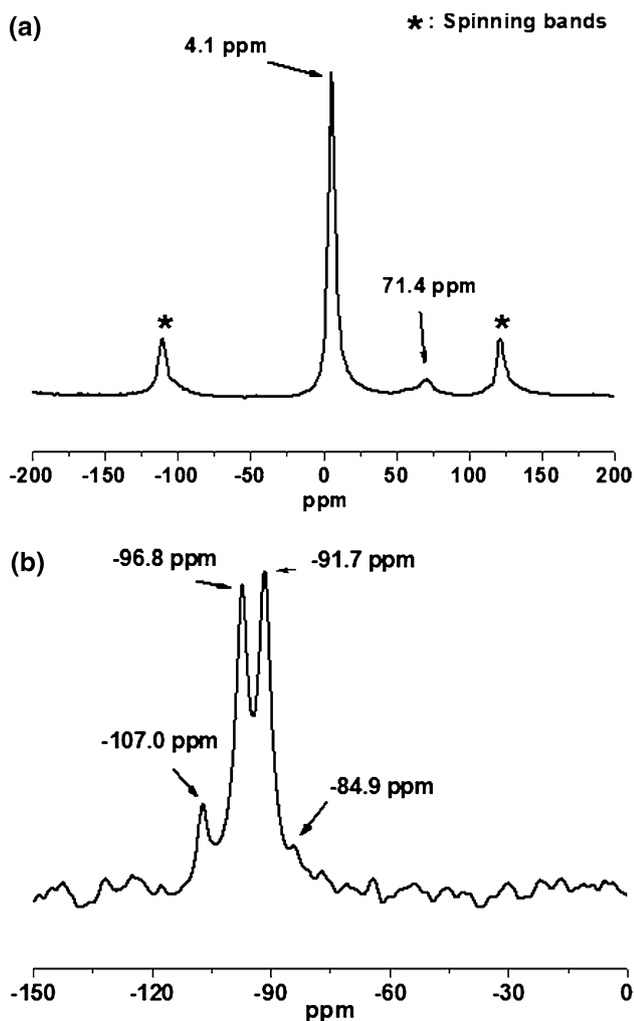


Fig. 6 ^{27}Al (a) and ^{29}Si (b) MAS-NMR spectra of Na^+ -PK7 $<2\ \mu\text{m}$ fraction

tetrahedral sheet of some Si^{4+} by Al^{3+} . The peak at $-107.0\ \text{ppm}$ corresponds to quartz impurity (Lippmaa et al. 1980) which remains in the samples even after purification in agreement with XRD analysis.

CEC and textural properties

The CEC, determined from the adsorption of cobalt hexamine (III) complex onto Na^+ -Pal according to the literature method (Mantin and Glaeser 1960), is found to be around $21.2\ \text{meq}/100\ \text{g}$. This value is of the same order of magnitude than for palygorskite from Oman ($19.2\ \text{meq}/100\ \text{g}$) and from Hawthorne, FL, USA ($19.5\ \text{meq}/100\ \text{g}$) (Al-Futaisi et al. 2007). Nevertheless, it is lower than those from other origins, for example, 25 , 42.5 , and $31\ \text{meq}/100\ \text{g}$ as reported by Serna et al. (1977), Blanco et al. (1989), and Chahi et al. (2002), respectively.

N_2 adsorption–desorption isotherms on raw clay and its fine fraction reveal that they are similar of type II, with

small hysteresis loops of type H3 according to the classification of the International Union of Pure and Applied Chemistry (IUPAC) (Sing et al. 1985) (Fig. 7). The shape of the isotherms denotes that both samples are mainly mesoporous solids, but also contain some micropores as clearly evidenced on the inset in Fig. 7 by the observation of a sharp increase at low pressure upon adsorption process. According to Cases et al. (1991), this microporosity is likely due to interfibers also revealed by SEM imaging (Fig. 3b), whereas channels could not be accessed to N_2 species and hence corresponding structural microporosity could not be detected owing to structure folding of palygorskite caused by prior outgassing treatment at $150\ ^\circ\text{C}$. The hysteresis of type H3 denotes that their particles form aggregates (Rouquerol et al. 2003). The BET specific surface area (S_{BET}) was determined from these isotherms, by applying BET equation for the relative pressure range $0.02 < p/p_0 < 0.33$. The microporous surface area (S_{mic}), external surface area (S_{ext}), and the micropore volume (V_{mic}) were obtained by the t-plot method (de Boer et al. 1966). Meanwhile, the mesopore volume (V_{mes}), the total pore volume (V_{T}), and pore size distribution were assessed according to the Barrett–Joyner–Halenda (BJH) method (Barrett et al. 1951). The overall results gathered in Table 1 show that, according to outgassing conditions (Cases et al. 1991; Jones and Galan 1988), the specific surface area of palygorskite sensibly varies from 100.4 to $116\ \text{m}^2/\text{g}$ when the outgassing temperature used is 150 and $140\ ^\circ\text{C}$, respectively. This value is still quite lower than the values reported in the literature, which are in the range 150 – $320\ \text{m}^2/\text{g}$ (Galán 1996).

Chemical analysis (ICP)

Chemical compositions of the dehydrated raw clay and its fine fraction indicate a huge decrease of CaO amount from about $37.6\ \%$ in the raw clay to around $0.3\ \%$ in the fine Na^+ -exchanged fraction (Table 2). Besides, the amount of potassium could be due to illite impurity existing in a minor amount associated to fibrous clay mineral as well as charge compensating cation in its channels. Illite was not detected by XRD analysis likely because of a proportion lower than the detection threshold and/or an overlapping of its characteristic (001) reflection ($\approx 10\ \text{\AA}$) (Holtzapffel 1985) with the very intense (110) basal reflection of palygorskite occurring at a close distance ($10.56\ \text{\AA}$).

At this stage, the partitioning of the amount of K among possible illite impurity and exchangeable compensating cation is not possible. The subtraction of the contribution of both sepiolite (assumed to be $5\ \%$) and quartz ($2\ \%$) from the composition of fine Na^+ -exchanged fraction (Na^+ -PK7) and taking into account the chemical mass balance relation (Hodgson and Dudeney 1984; Engler and

Iyengar 1987; Bruand and Prost 1988; Bouabid and Badraoui 1996) yields to the elemental composition of dehydrated palygorskite (Na^+ -Pal) and the number of cations per 21 atoms (Table 2). The structural formula of dehydrated palygorskite determined by neglecting the contribution of K is approximately found to be $(\text{Si}_{7.92}\text{Al}_{0.08})(\text{Mg}_{2.15}\text{Al}_{1.4}\text{Fe}_{0.4}\text{Ti}_{0.05}\square_1)(\text{Ca}_{0.03}\text{Na}_{0.08})\text{O}_{21}$. This formula shows that the tetrahedral occupation of Al represents 5 % in good agreement with the proportion beforehand determined from the ratio $\text{Al}^{\text{IV}}/(\text{Al}^{\text{IV}} + \text{Al}^{\text{VI}})$ of ^{27}Al MAS-NMR peaks intensities. With respect to the ideal formula $\text{Si}_8\text{Mg}_5\text{O}_{20}(\text{OH})_2(\text{OH}_2)_{4,4}(\text{H}_2\text{O})$ proposed by Bradley (1940) for a pure trioctahedral palygorskite, the determined formula of the dehydrated palygorskite from Marrakech High Atlas shows excess of negative charges on tetrahedral and octahedral sheets of about -0.08 and -0.1

e/half unit, respectively. They are compensated by a lower positive charge ($+0.14$ e/half unit) due to Ca^{2+} and Na^+ interlayer cations. This positive charge deficit could be completed by introducing 0.04 of exchangeable compensating K^+ ions to ensure the electrical neutrality of the overall formula. This deduced amount of K^+ corresponds to 0.23wt% of K_2O which is of the same order of magnitude than those reported to be usually contained in several palygorskites from other areas (Chahi et al. 2002; Garcia-Romero and Suárez 2010). Hence, the most appropriate chemical formula per half unit cell of the dehydrated palygorskite investigated herein, deduced on the basis of 21 oxygens, could be $(\text{Si}_{7.92}\text{Al}_{0.08})(\text{Mg}_{2.15}\text{Al}_{1.4}\text{Fe}_{0.4}\text{Ti}_{0.05}\square_1)(\text{Ca}_{0.03}\text{Na}_{0.08}\text{K}_{0.04})\text{O}_{21}$.

The structural formula of the hydrated palygorskite, calculated on the basis of 26 oxygens by taking into account the experimental amounts of different kinds of water determined above by TG (Caillère and Hénin 1961), is $(\text{Si}_{7.97}\text{Al}_{0.03})(\text{Mg}_{2.17}\text{Al}_{1.46}\text{Fe}_{0.40}\text{Ti}_{0.05})(\text{Ca}_{0.03}\text{Na}_{0.07}\text{K}_{0.03})\text{O}_{20.18}(\text{OH})_{1.94}(\text{OH}_2)_{3.88}\cdot 2.43 \text{ H}_2\text{O}$. This crystal-chemical formula shows that the octahedral sheet composition exhibits a predominant dioctahedral character with divalent to trivalent cation ratio $R(\text{Al}^{3+} + \text{Fe}^{3+})/R(\text{Mg}^{2+})$ around 0.84. Besides, the same formula evidences the above commented deficiency of zeolitic H_2O (2.43 molecules per crystal-chemical formula instead of 4), as inferred by TGA data. The M_2 sites linked to structural OH seem to be predominantly occupied at 90 % by trivalent cations, that is, Al^{3+} (78 %) and Fe^{3+} (22 %), at 7.5 % by divalent cations Mg^{2+} and at 2.5 % by tetravalent cations Ti^{4+} .

On the basis of all octahedral cations occupation percentages determined herein on the one hand and knowing that palygorskite octahedral sheet contains 10 charges per half-cell on the other hand, a qualitative octahedral cation distribution for the palygorskite from Marrakech High

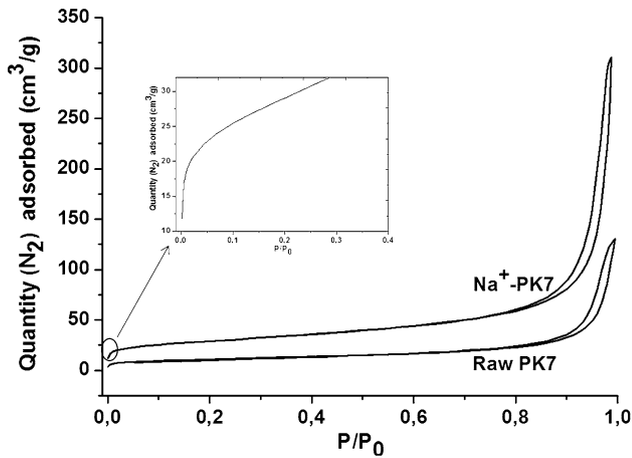


Fig. 7 Adsorption–desorption (N_2) isotherms of the raw clay (PK7) and the separated fine fraction (Na^+ -PK7)

Table 1 BET specific surface area (S_{BET}), total pore volume (V_{T}), microporous volume (V_{mic}), mesoporous volume (V_{mes}), macroporous volume (V_{mac}), micropore surface area (S_{mic}), and external surface area (S_{ext}) of the raw clay (PK7) and its purified $<2 \mu\text{m}$ fraction (Na^+ -PK7)

Clay	S_{BET} (m^2/g)	Total pore volume, V_{T} (cm^3/g)	Micropores volume, V_{mic} (cm^3/g)	Mesopores volume, V_{mes} (cm^3/g)	Macropores volume, V_{mac} (cm^3/g)	Micropores surface area, S_{mic} (m^2/g)	External surface area, S_{ext} (m^2/g)
Raw PK7	38.4	0.1937 ^d	0.0004 ^e	0.0693 ^e	0.1224 ^f	4.2	34.2 ^g
Na^+ -PK7	100.4 ^a 112.7 ^b 116.7 ^c	0.4581 ^d	0.0020 ^e	0.1632 ^e	0.2927 ^f	12.2 12 22.9	88.2 ^g 100.7 ^g 93.8 ^g

^a S_{BET} determined by outgassing sample at 150 °C for 24 h

^b S_{BET} determined by outgassing sample at 150 °C for 12 h

^c S_{BET} determined by outgassing sample at 140 °C for 9 h

^d Determined from BJH adsorption cumulative pore volume of pores having diameters in the range 17.0–3000.0 Å

^e Determined from BJH adsorption cumulative pore volume data for pores having diameters in the range 20.0–500.0 Å

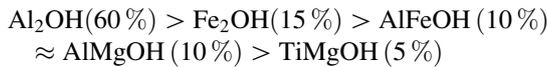
^f Determined from BJH adsorption cumulative pore volume data for pores having diameters above 500.0 Å

^g Determined from the t -curve (adsorbed volume, V_{ads} vs. the thickness t)

Table 2 Chemical compositions of the raw clay (PK7) and its <2 μm fine fraction (Na^+ -PK7)

Samples Oxides	Raw clay (PK7) Wt %	Na^+ -PK7 (0.2 N) Wt %	Na^+ Pal Wt %	Cations number/half-cell
SiO_2	33.00	66.26	64.29	7.92
Al_2O_3	8.35	9.83	10.23	1.48
Fe_2O_3	4.00	4.18	4.32	0.40
MgO	11.76	2.51	11.74	2.15
CaO	37.62	0.27	0.26	0.03
Na_2O	0.65	0.32	0.32	0.08
K_2O	1.37	2.01	2.01	–
TiO_2	0.28	0.60	0.60	0.05
LOI	2.97	4.04	6.23	–
Total	100	100	100	

Atlas could be illustrated by considering 10 unit cells as in Fig. 8. In this distribution, the rate of hydroxyl groups linked to M_2 sites seems to increase as a function of octahedral cations as follows:



The CEC corresponding to the compensating cations charge ξ (+0.16 e/half unit cell) exhibited by the structural formula above can be calculated from the following Eq. (1) (Besson et al. 1990; Mermut and Lagaly 2001):

$$\text{CEC} = 10^5 \times \xi / M \quad (1)$$

where M designates the molar mass. The calculated value about 20 meq/100 g is quite similar to the experimental value measured from cobalt hexamine adsorption.

Octahedral sheet composition (IR)

As beforehand revealed by XRD analysis (Fig. 4a) and thermal analysis (Fig. 5), the MIR spectra of the raw clay and its fine Na^+ -exchanged fraction confirm the entire removal of carbonates impurities largely present in the raw clay upon the purification treatment as evidenced by the disappearance of corresponding bands at 1,440, 870, and 714 cm^{-1} (Fig. 9a). Except for these differences, the MIR spectra of the two samples reveal similar absorption bands which are better resolved by superposing the corresponding second-derivative spectra (Fig. 9a).

The octahedral sheet composition deduced above is further strongly supported by IR spectroscopy in middle and near frequencies regions which generally gives evidence for strong vibration bands of Al_2OH and relatively small peaks of Fe_2OH and AlFeOH . Indeed, the bands at 3,618, 3,555, and 3,580 cm^{-1} are due to the stretching OH vibrations in Al_2OH and/or of Mg-coordinated OH_2 ,

Fe_2OH , and AlFeOH and/or AlMgOH species, respectively (Serna et al. 1977; Blanco et al. 1989; Gionis et al. 2006, 2007). Nevertheless, the contribution of AlMgOH seems to be very minor since the above determined structural formula shows very low amount of Mg in central positions (M_2 site) of octahedral sheet. Their corresponding deformation vibrations' bands are neatly observed at 910, 800, and 780 cm^{-1} , respectively (Gionis et al. 2006, 2007). These assignments are further confirmed in the second-derivative NIR spectrum of the fine Na^+ -exchanged fraction by the presence of a well-resolved triplet of bands at 7,059, 7,000, and 6,926 cm^{-1} (Fig. 9b) corresponding to the first overtones of the O–H stretching modes ($2\nu_{\text{OH}}$) of Al_2OH , AlFeOH , and Fe_2OH entities, respectively (Gionis et al. 2006, 2007).

Furthermore, their corresponding stretching–deformation combination vibrations are also well detected in the second-derivative NIR spectrum at 4,498, 4,431, and 4,340 cm^{-1} , respectively (Fig. 9b). The observation of an additional well-defined peak at 3,684 cm^{-1} (Fig. 9a) is worth noting. It likely corresponds to the stretching vibration of OH in Mg_3OH species (Chahi et al. 2002). The corresponding deformation band is observed at 680 cm^{-1} , while its corresponding first overtone of the O–H stretching vibration is clearly observed at 7,215 cm^{-1} in the second-derivative NIR spectrum (Fig. 9b). A broad peak centered at around 1,650 cm^{-1} is observed whose second derivative reveals that it is made up of two well-resolved peaks at 1,640 and 1,660 cm^{-1} (Fig. 9a), assigned to coordinated and zeolitic H_2O present in palygorskite according to Gionis et al. (2006) and Gionis et al. (2007). A third band at 1,675 cm^{-1} , ascribed by these authors to physisorbed water, is not observed in the samples investigated herein likely because of their preparation (storage and drying) prior analysis. However, the corresponding stretching vibrations bands that are more sensitive are clearly observed at 3,270 and 3,430 cm^{-1} in the two samples.

Furthermore, the second-derivative NIR spectrum of the Na^+ -PK7 sample also reveals the presence of a main peak at 5,210 cm^{-1} and a shoulder at 5,045 cm^{-1} due to combination modes of the three types of H_2O (physisorbed, zeolitic, and coordinated to edge cations) characteristic of palygorskite. The bands at 1200 and 646 cm^{-1} are fingerprints of fibrous clay minerals as palygorskite and sepiolite. They correspond to the asymmetric and symmetric stretching of the Si–O–Si bridges linking the aluminosilicate slabs of these fibrous clay minerals (Gionis et al. 2006). The corresponding deformation vibration band clearly appears at 428 cm^{-1} . The band at 1,093 cm^{-1} could be ascribed to stretching vibrations of Al–O in octahedral sheet (Blanco et al. 1989). The absorption bands at 1,028 and 985 cm^{-1} are characteristic of antisymmetric

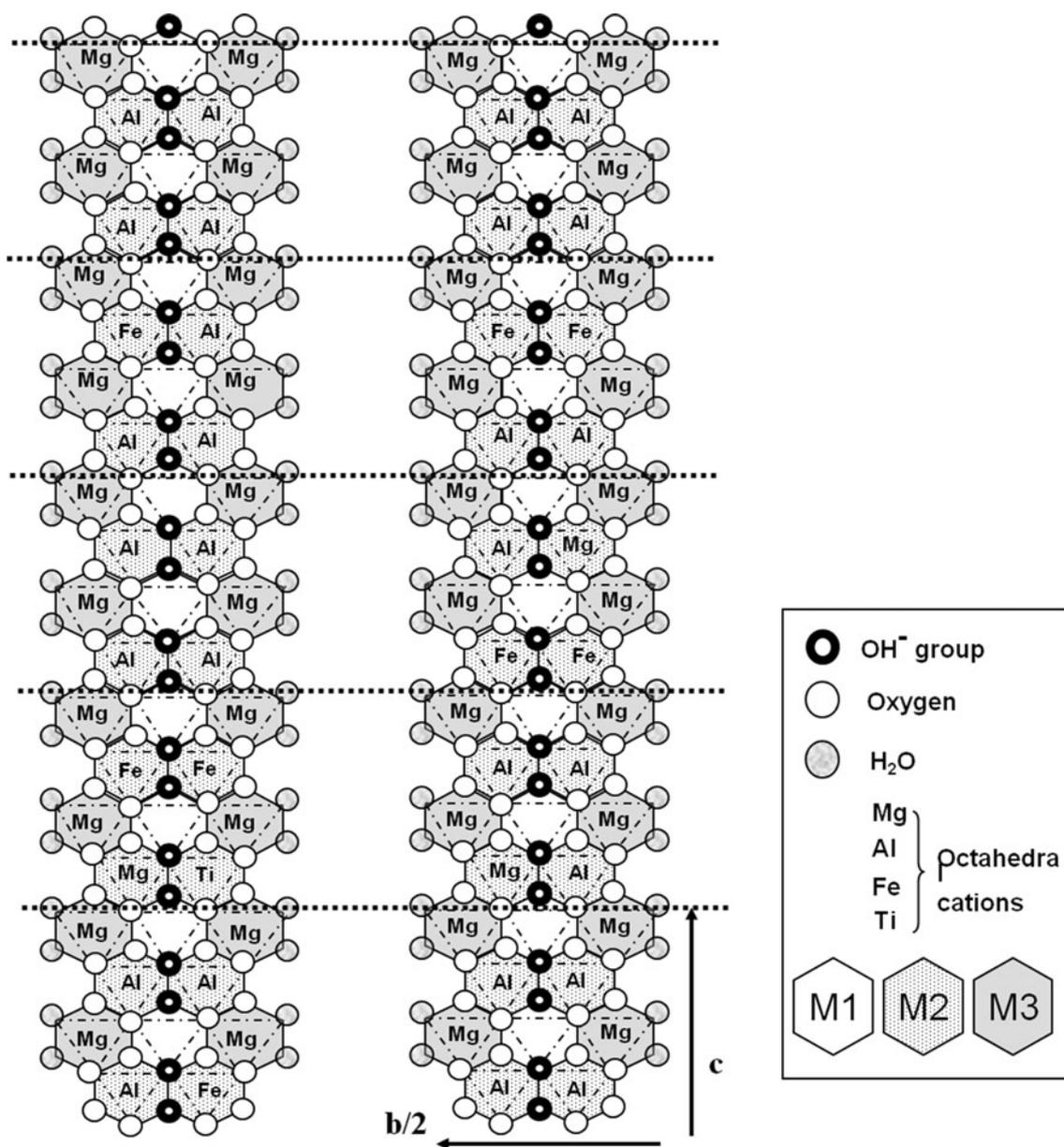


Fig. 8 Schematic distribution of cations in the octahedral sheet of ten units of palygorskite from Marrakech High Atlas [projection on the (100) plane] showing that the 20 most inner M_1 sites were vacant and

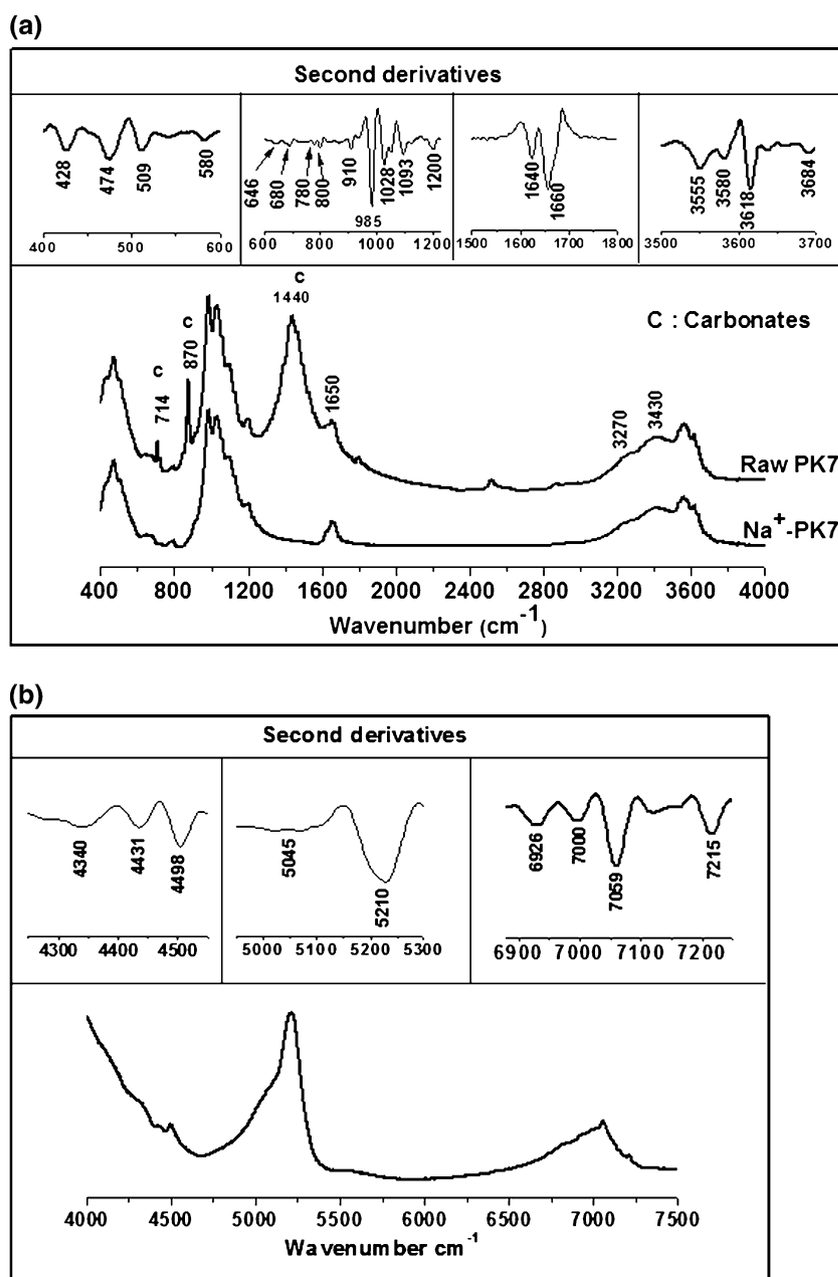
the following 40 M_2 sites were shared between 28 Al^{3+} , 8 Fe^{3+} , 3 Mg^{2+} , and 1 Ti^{4+} , while the 40 M_3 sites at the edges were all occupied par Mg^{2+}

vibrations of Si–O and Al–O in tetrahedral sheet of palygorskite (Augsburger et al. 1998; Suárez and García-Romero 2006). The corresponding deformation vibration bands are clearly observed at 474 and 580 cm^{-1} for Si–O bonds. The peak at 509 cm^{-1} is also observed by Suárez and García-Romero (2006) for Al-rich palygorskites; it is ascribed to vibrations of Al–O–Si in tetrahedral sheet. These results denote the substitution of Al^{3+} for Si^{4+} in tetrahedral sheet as also revealed by NMR analysis (Fig. 6) and the determined structural formula.

Conclusions

This study demonstrates that the fine fraction (<2 μm) isolated from clay, picked up in Marrakech High Atlas (Morocco), is rich in palygorskite (95 %) along with minor amount of sepiolite (5 %). The crystal-chemistry formula of this palygorskite has been determined as: $(Si_{7.97}Al_{0.03})(Mg_{2.17}Al_{1.46}Fe_{0.40}Ti_{0.05})(Ca_{0.03}Na_{0.07}K_{0.03})O_{20.18}(OH)_{1.94}(OH_2)_{3.88} \cdot 2.43 H_2O$. This shows a deficiency in zeolitic H_2O (5 wt%) which seems to be more pronounced than for

Fig. 9 Mid-infrared spectra (a) and near-infrared spectra (b) of clay samples before (PK7) and after the purification treatment (Na⁺-PK7)



palygorskite from Montestrutto, Italy (Giustetto and Compagnoni 2011: 6 wt%). Besides, the ratio (Al³⁺+Fe³⁺)/Mg²⁺ of about 0.84, inferred from the chemical composition of octahedral sheet, indicates a pronounced dioctahedral character of this palygorskite in contrast with the palygorskite from phosphorite deposits of Gantour (Morocco) exhibiting a mixture of di- and trioctahedral property (Chahi et al. 2002). This is further supported by the observation of intense fundamental OH stretching modes in MIR spectra as well as their corresponding first overtones in NIR spectra arising from mainly Al₂OH units along with those relatively less intense of Fe₂OH and AlFeOH groups. On the basis of these premises, a scheme (Fig. 8), elucidating the octahedral

sheet composition of palygorskite from Marrakech High Atlas, was proposed. Its functional features, in terms of cation exchange capacity, specific surface area, and total pore volume, were also assessed to be about 21.2 meq/100 g, 116 m²/g, and 0.46 cm³/g, respectively.

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