CHARACTERIZATION OF ALTERED MICA FROM SOKLI, NORTHERN FINLAND

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Abstract—Vermiculite is a mineral with many potential uses in various industrial areas, such as in insulation, horticulture, and environmental applications. The regolith of the Sokli carbonatite intrusion in northern Finland consists of weathered micas which reportedly contain vermiculite. The aim of the present study was to characterize the weathered mica in order to determine if the weathered regolith contains vermiculite. If so, the value of the apatite-rich Sokli intrusion may increase, because the vermiculite could be classified as an ore in Sokli. For the characterization, mica fractions were investigated using electron probe microanalysis (EPMA), X-ray diffraction (XRD), and thermogravimetric analysis (TGA). The EPMA results show that the potassium (K2O) content of the mica investigated varies between 6.20 and 0.31 wt.%. The XRD results provided evidence that the interlayer distances in the mica vary between 10.1 and 14.7 Å. The TGA shows that the dehydration of the samples varies between 6 and 12 wt.% for temperatures up to 170°C. Dehydroxylation takes place in three steps, as is characteristic for vermiculite. The results from this study revealed that mica in the weathered parts of the Sokli carbonatite is mostly vermiculite, and that the vermiculite was formed by the weathering of phlogopite. These results can be used to determine whether vermiculite is a possible future commodity at the Sokli carbonatite complex.

Keywords—EPMA · Phlogopite Alteration · Sokli Carbonatite · TGA · Vermiculite · Vermiculitization · XRD

INTRODUCTION

The demand for vermiculite has increased recently. The physical and chemical properties of natural vermiculite, such as low density, high cation exchange capacity, and high adsorption ability (Bergaya & Lagaly 2013), are the reasons for using vermiculite in building materials, the chemical industry, and in various applications in environmental protection and agriculture (Huo et al. 2012). Among these uses are as non-flammable packing material, water purification by ion exchange (Rama et al. 2019), and thermal insulation (Bergaya & Lagaly 2013). Exfoliated vermiculite is prepared by heating to >300°C and is used in many commercial applications, such as heat-insulating materials or soil improvers (Suvorov & Skurikhin 2003). Vermiculite is not mined in Finland even though it has been reported to exist at the Sokli carbonatite intrusion (Vartiainen 1974; Kresten & Berggren 1978; O’Brien & Hyvönen 2015).

The Sokli complex is the western-most intrusion of the Kola alkaline province. It is the only part of the province located outside Russia, and is situated 60 km NNW of the Kovdor complex (Krasnova 2001; Lee et al. 2003). Kovdor is known for its rich quarries from which magnetite, apatite, baddeleyite, phlogopite, and vermiculite are excavated. The Sokli massif (and the Kovdor complex) consists principally of intrusive carbonatite stock with a fenite aureole. In addition to the carbonatites, the massif is composed of ultramafic rocks (dunite and pyroxenite) and multiple stages of phoscorites (forsterite-magnetite-apatite-calcite rocks). Phlogopite is a common silicate in the Sokli carbonatite (Lee et al. 2003) and vermiculite has reportedly been found both in the carbonatite stock and in the regolith derived from weathering of the phoscorite (Kresten & Berggren 1978). The phoscorite layer in Sokli has a commercial value for the production of phosphorus fertilizers. Vermiculite from the Kovdor regolith has been excavated for commercial use, and Rama et al. (2019) reported a comparison between Sokli and Kovdor vermiculite.

Vermiculite (Mg,Fe,Al)3(Al,Si)4O10(OH)2·4(H2O) is a mixed-layer phyllosilicate of the hydromica group, and is formed by hydration and other secondary changes of micas, particularly phlogopite (K(Mg,Al)sI3O10(F,OH)2) and biotite (K(Mg,Fe)sI3O10(F,OH)2) (Ghabru et al. 1989; Deer et al. 1992). According to Roy & Romo (1957) and Komarneni & Roy (1981), vermiculite is a product of mica alteration by dilute, low-alkali solutions. Alternation of different layers along the c* direction can create either ordered or regular mixed-layer structures (Brigatti et al. 2006). Mixed-layer clay minerals with any appreciable degree of randomness show an irregular series of basal reflections in XRD patterns. Mixed-layer biotite-vermiculites are, in most cases, derived from the degradation of pre-existing clay minerals. Previous studies indicated that tri-octahedral biotite could weather to vermiculite under acid conditions, and to vermiculite + montmorillonite under neutral and alkaline conditions (Latimer 1952). The oxidation of Fe(II) to Fe(III) under alkaline conditions causes a large decrease of the clay mineral surface charge and an expansion of the crystal lattice. Based on the analysis of vermiculites and ‘hydrobiotites,’ Foster (1963) concluded that the composition, charge relation, and cation.
exchange capacity in these vermiculites can be derived from phlogopite or magnesian biotite by the replacement of K$^+$ by Mg$^{2+}$. The term ‘hydrobiotite’ was interpreted as regular 1:1 interstratification of biotite (or interlayer-deficient biotite) and vermiculite (Rieder et al. 1998; Valašková et al. 2018).

The silicate layers are negatively charged, and electrical neutrality is preserved by the presence of cations between the layers. These interlayer cations are usually Mg$^{2+}$ or Ca$^{2+}$, associated with double layers of water molecules. The interlayer ions are exchangeable and can be replaced by various monovalent or divalent cations (e.g. Li$^+$, Na$^+$, K$^+$, Cs$^+$, NH$_4^+$, Ni$^{2+}$). Pure vermiculite has a cation exchange capacity (CEC) of 1.2–2.4 meq/g (Deer et al. 1992). In addition to the layered structure and large CEC, the third important feature of the vermiculite structure is the variable amount of water in the interlayer space (Tunega & Lischka 2003). The degree of hydration of interlayer cations is a function of the partial vapor pressure of water, the cations, and the charge density in the layers (Whittingham 1989). Due to hydration, the structure of vermiculite can be modified with heat (Kresten & Berggren 1978; Marcos et al. 2009; Mouzdahir et al. 2009).

The occurrence of vermiculite in Sokli has been reported by several authors (Kresten & Berggren 1978; O’Brien & Hyvönen 2015), though there has been no detailed research to confirm the vermiculite findings and the origin of the vermiculate. Prior to commercial utilization, assessment of the quality and estimates of vermiculite reserves are needed. The purpose of the present study was to test the hypothesis that mica flakes from Sokli are mostly vermiculite by performing detailed mineralogical analyses of mica flakes from the weathered surface of the Sokli massif, using a battery of methods, including EPMA, TGA and XRD.

GEOLOGICAL BACKGROUND AND FIELD RELATIONSHIPS

The Sokli carbonatite complex is located in northeastern Finland, with a center point near 67° 48’12.2” north latitude and 29° 19’2.3” east longitude (O’Brien & Hyvönen 2015). The complex was discovered in 1967 in an airborne geophysical survey by the Rautaruukki Company (Paarma 1970). The Sokli carbonatite complex is slightly oval in outline, with a major axis oriented northwest-southeast of ~6.2 km and a minor axis of 4.8 km, covering an area of 20 km$^2$ (Lee et al. 2003; O’Brien & Hyvönen 2015). Deep seismic sounding data have revealed a conical shape that tapers slightly with increasing depth (Vartiainen 1980). The intrusion consists mainly of calcite carbonatite surrounded by a fertilized aureole (Paarma 1970). The Rb/Sr age of the Sokli carbonatite is 365 ± 3 Ma (Mäkelä & Vartiainen 1978; Kramm et al. 1993). The complex is divided into five geological units (Fig. 1): (1) magmatic core; (2) metasomatic carbonatite surrounding the core; (3) phoscorites and metasomatic phoscorites occurring as large blocks within the carbonatite; (4) transitional zone between the metacarbonatites and fenites; and (5) fenite aureole (Vartiainen 1980; Vartiainen & Vuotovesi 1980; O’Brien et al. 2005).

The magmatic carbonatites of the inner core developed successively in various stages, first by direct crystallization of a carbonatite magma, and later by hydrothermal processes (Vartiainen 1980). The metacarbonatites dominate an area quite clearly outlined against the central core, whereas the contacts with the surrounding transition zone are gradual. The main rock type of the metacarbonatites is metasomatic carbonatite, which was formed by metasomatic replacement of an unidentified lithology by the introduction of CO$_2$-rich fluids derived from the carbonatite magma. An arcuate, broken zone of metaphoscorite is located between the metacarbonatite area and the transition zone. The zone was composed initially of ultrabasic rocks comprising magnetite-bearing dunites (partly of pegmatoidal grain size), pyroxenites, and amphibole rocks (Vartiainen 1980). The metaphoscorites are considered as carbonatized alteration products of ultramafites (Vartiainen & Woolley 1976). During multiple metasomatic alteration processes, aegirine, amphiboles, micas, apatite, carbonates, sulfides, and zircons were accommodated in the ultramafites. Within the magmatic core, where metaphoscorites were trapped during the intrusion of the coreplug, the variable effects of magmatic carbonatites complicated still further the development of metaphoscorites (Vartiainen 1980). The transition zone of the metasomatisms forms a girdle between the fenite aureole and the metacarbonatite area. The metasomatized zone is composed of fragments of fenites, pyroxenites, omphaloids, mica-amphibole rock, and mica rocks embedded in metalamcosovites (Vartiainen 1980). The fenite aureole is divided into inner syenitic and outer, low-grade fenite areas. The syenitic fenites are completely recrystallized massive rocks with fractures filled by carbonate, aegirine, and sodium amphiboles (Vartiainen & Paarma 1979; Vartiainen 1980).

The Sokli area is covered by weathered bedrock of variable thickness. The most important mineralogical weathering processes are the partial dissolution of carbonates, local replacement of phlogopite by vermiculite, intense alteration of olivine, the partial replacement of magnetite by hematite, alteration of pyrochlore, and the total removal of sulfides (O’Brien & Hyvönen 2015). The intrusion is capped by a 26-meter-thick regolith that constitutes the main phosphate ore. The phosphorus ore developed from the carbonatite that underwent complex weathering, leaching, recrystallization, and lithification processes (O’Brien & Hyvönen 2015). The P$_2$O$_5$ content in the phosphorus ore ranges from 4 to 5 wt.% to up to 30 wt.%. The consulting and engineering firm Pöyry Environment Oy (2009) estimated the regolith ore reserves to be ~75 Mt by limiting reserves to areas under the thinnest overburden in the Kaulusrova area, where the grade of P$_2$O$_5$ is 5.6 wt.%. In addition to phosphate ore, the deposit contains Nb, Ta, Zr, and some U. According to Vartiainen (1974), the regolith contains up to 10 wt.% mica and the vermiculite content is up to 6 wt.% in the area.

MATERIALS AND METHODS

Sampling and sample preparation

Soil samples were collected from the regolith altered from carbonatite and phoscorite in the Kaulusrova deposit of the Sokli complex (Sokli9 and Sokli10; Fig. 1). The soil samples
consist mainly of mica and phoscorite, calcite, carbonatite, magnetite, pyroxenites, and amphibole. The soil samples contained ~50% of mica flakes with a grain size varying from ~1 mm to 10 cm. To avoid possible structural damage, the mica flakes were separated from the soil by hand and without any chemical pretreatments or dispersants.

The mica flakes were characterized by EPMA, XRD, and TGA. For XRD and EPMA analyses, mica flakes with a surface size of 11 mm×12 mm and a 3–6 mm thickness from sample Sokli10 were mounted in epoxy and polished (Fig. 2). A Sokli9 sample was used for TGA analysis. This sample was divided into three fractions: a solid sample, Sokli1, with a grain size ~3 cm across and two ground samples, Sokli7 and Sokli8, with a grain size of between 0.075 and 0.125 mm.

**Analytical methods**

The elemental chemical composition of the mica was obtained from wavelength dispersive electron probe microanalysis (WD-EPMA) (JEOL: JXA-8600 superprobe, Tokyo, Japan) at Top Analytica, Turku, Finland, using an accelerating voltage of 15 kV and a beam current of 22 nA. Elemental concentrations were calibrated against natural and synthetic minerals, or pure metals. Data were acquired and ZAF corrections were performed using XMAS software (SMAX 1993).

The XRD measurements were performed using a Bruker D8 Discover (Karlsruhe, Germany) instrument (Laboratory of Physical Chemistry, Åbo Akademi University, Turku, Finland) using a Cu Kα X-ray source (λ= 1.5406 Å) and a scintillation detector. The beam was focused using a 0.5 mm collimator. The various measurement points on the sample were scanned in the range 3–75°2θ with an increment of 0.04°2θ/0.5 s. The measurement points were taken for each 1.5 mm on the surface of the sample, and for each 1.0 mm on the cross section.

The thermal response of the mica was studied using a thermogravimetric analyzer (TGA) and differential scanning calorimeter (DSC) - TA Instruments SDT Q600 (New Castle, Delaware, USA) between 25 and 1200°C in flowing air (flow rate: 100 cm³/min⁻¹) (Laboratory of Materials Chemistry and Chemical Analysis, University of Turku, Turku, Finland). Samples of 10 mg each were heated at a rate of 5°C/min. The sample pan
material was aluminum oxide and an empty aluminum oxide pan was used as a reference. The specific enthalpies of the DSC signals were calculated by integration.

RESULTS

XRD results

X-ray diffraction was used to investigate the layer structure and to estimate qualitatively what phases are present in sample Sokli10. The XRD pattern (Fig. 3) clearly displays a main reflection at 6.04°2θ. This strong peak is typical of Mg-vermiculite and can be attributed to the 002 plane, corresponding to an interlayer distance of 14.7 Å (Al-Ani & Saranpää 2008; Barabaszová & Valášková 2013; da Silva et al. 2013). The peak at position 8.73°2θ corresponds to d = 10.1 Å and indicates a small amount of phlogopite (K\text{Mg}_3\text{Al}_3\text{Si}_5\text{O}_{10}(\text{F,OH})_2), as phlogopite has a basal 002 spacing of 10.0–10.5 Å (Bailey 1980; Chmielarz et al. 2010). Other peaks were detected at 12.07°2θ (d = 7.32 Å), 18.02°2θ (d = 4.88 Å), 24.21°2θ (d = 3.66 Å), 30.39°2θ (d = 2.93 Å), 43.07°2θ (d = 2.10 Å), and 63.22°2θ (d = 1.47 Å). This XRD pattern is characteristic of vermiculite, although d spacings of vermiculites and other expandable phyllosilicates can vary slightly with changes in relative humidity. Both the intensities and the shapes of the reflections are typical of vermiculite data obtained from the literature (Bailey 1980; Vali & Hesse 1992; Mališ & Kříšková 2005; Campos et al. 2009; Hongo et al. 2012; Valášková et al. 2013; Padilla-Ortego et al. 2014). All of the XRD patterns of the vermiculite sample, measured from several points perpendicular to the surface, imply an overall well ordered layered structure. Some small impurities were detected at two positions in the sample. The structure is almost perfectly aligned, as no diffraction peaks could be detected when irradiating the cross-section.

Figure 3. XRD pattern of the Sokli10 sample. Crystal plane indices are noted in parentheses and values in Å refer to the spacing. P = phlogopite peak

Figure 4. Thermogravimetry (TG) and Differential Scanning Calorimetry (DSC) curves for vermiculite from Sokli
The XRD patterns of the mica analyzed are clearly similar to those typical of vermiculite with a main reflection 14.7 Å (002). A weak reflection at 10.1 Å (002) indicates interstratification of the mixed-layer structure between phlogopite and vermiculite (Brigatti et al. 2006; Valsásková et al. 2018). Interstratification of phlogopite layers and vermiculite layers are visible in a single grain in Figs 7a and 7b.

**Temperature dependence results**

Thermogravimetry and DSC results are presented to assess the behavior of Sokli vermiculites during dehydration. The temperatures of dehydration reactions of vermiculite vary depending on the relative humidity and the degree of interstratifications with other phyllosilicates, i.e. biotite or phlogopite/vermiculite interstratifications. Due to hydration, the structure of vermiculite can be tailored by thermal treatment.

The dehydration of vermiculite upon heating usually takes place in three steps (Földvári 2011): (1) below and around 100°C, the water adsorbed on the surface is lost; (2) around 200°C, the water between the layers evaporates; and (3) at 250–300°C the water bound to the exchangeable cation escapes.

The TG curves (Fig. 4) show dehydration between room temperature and 200°C only. Two steps of dehydration amounting to 6 wt.% (Sokli7) and 12 wt.% (Sokli8) correspond to loss of the water bound to the exchangeable cation in place of two hydroxide groups. The first dehydration step differs in magnitude between the two samples, but the second step corresponds to ~3 wt.% for both samples. After this dehydration, the TG curve shows a plateau due to the formation of (Mg,Ca,Fe,Al)((Al,Si)4O10)(OH)2. For the bulk sample (Sokli1), no clear plateau in the TG curve is seen around 300°C (Földvári 2011). For each sample, the dehydroxylation takes place in three steps (Table 2), but whether these involve endo- or exothermic changes, if any, is difficult to say based on the DSC data. For the ground samples (Sokli7 and Sokli8), the dehydroxylation corresponds to a total mass loss of 8%, whereas the weight loss from the bulk sample was ~9%. The endothermal release of heat at 830°C for samples Sokli7 and Sokli8, and at 895°C for sample Sokli1 (Fig. 4), is a characteristic feature of vermiculite and probably represents the recrystallization to enstatite (Balek et al. 2007; Földvári 2011).

Comparing the mass losses associated with dehydration and dehydroxylation, and assuming an initial composition of (Mg,Ca,Fe,Al)((Al,Si)4O10)(OH)2·nH2O, leads to the conclusion that sample Sokli7 is a dihydrate and sample Sokli8 is a trihydrate, whereas sample Sokli1 consists of a mixture of the two. Because sample Sokli1 was not ground, thermal conductivity is lower than that of the ground samples. The dehydration for sample Sokli1 may already have occurred at temperatures up to well above 200°C and the dehydroxylation started before the dehydration was complete, making quantification of the decomposition process more difficult.

X-ray powder diffraction patterns were obtained for all three samples in as-received form as well as after heating for 1 h at 600°C in air. The patterns of the natural vermiculite samples are dominated by the strong (001) reflection at 6°2θ (Fig. 5). The XRD data confirmed that samples Sokli7 and Sokli8 are very similar in composition, which is in agreement with their very similar thermal decomposition. On the other hand, sample Sokli1 is the purest vermiculite of these samples. The XRD data explained why sample Sokli1 behaves clearly differently in the TG measurements.

The XRD patterns recorded after heating the powders for 1 h at 600°C confirmed the expansion of the vermiculite structure, as evidenced by the shift of the reflections to smaller angles (Figs 5 and 6). Also, the loss of some reflections indicated the decomposition of impurities. The expansion is best observed from the shifting of the (001) reflections (Table 3). The dominant (002) reflection of each sample at 5.9°2θ shifted to angles that are smaller than the low-angle

### Table 1 Characteristics of dehydration for Sokli vermiculite

| Sample | First loss of water (25–75°C) | Second loss of water (75–170°C) | Total |
|--------|--------------------------------|---------------------------------|-------|
|        | Loss (%) | ΔH (J g⁻¹) | Loss (%) | ΔH (J g⁻¹) | Loss (%) |
| Sokli1 | 8        | 91.3      | 1        | 2.7        | 9       |
| Sokli7 | 3        | 19.9      | 3        | 65.1       | 6       |
| Sokli8 | 9        | 144.2     | 3        | 64.7       | 12      |

### Table 2 Characteristics of dehydroxylation

| Sample | First step | Second step | Third step |
|--------|------------|-------------|------------|
|        | Loss (%) | T (°C) | Loss (%) | T (°C) | Loss (%) | T (°C) |
| Sokli1 | 5        | 155–690  | 1        | 690–765 | 3        | 765–940 |
| Sokli7 | 1        | 420–520  | 3        | 520–685 | 4        | 685–915 |
| Sokli8 | 1        | 410–515  | 3        | 515–690 | 4        | 690–930 |
limit of the detector (4.0°2θ), and the expansion had to be studied from the shift in (0010) and (0012) reflections.

**EPMA results**

Back-scattered electron (BSE) images revealed variations in the chemical composition of mica sample Sokli10. Point analyses showed that brighter areas contained more potassium than the darker areas (Fig. 7). The interlayer distance in the sample varied from 10.1 to 14.7 Å, where 10.1 Å was similar to phlogopite and 14.7 Å was similar to Mg-vermiculite (see the XRD results section).

Quantitative chemical analyses were performed on mica sample Sokli10 (Table 1). The amounts of SiO₂, Al₂O₃, MnO, and TiO₂ were relatively constant. MgO and Fe₂O₃ composition varied between 25.05 and 17.52 wt.% and 4.17–6.94 wt.%, respectively. The vermiculitization process can be seen in the XRD results as an increase in interlayer space. The areas with the thinnest interlayer have the largest K₂O content, while the layers with the thickest interlayer are depleted in K₂O. Additionally, the areas with the largest K₂O content also have the largest total, while K₂O-depleted micas have the smallest total. The decrease in analytical total may indicate an increase in the non-measured water content in the interlayer of the mica. The decrease in K₂O also represents the vermiculitization process. As K₂O (and Na₂O) decreases, CaO is simultaneously enriched from 0.02 wt.% to 1.88 wt.%. Due to this variation in analyzed points, the vermiculitization process was apparent in this sample.

![Figure 5](https://doi.org/10.1007/s42860-019-00041-0) Effect of heating on the XRD patterns

![Figure 6](https://doi.org/10.1007/s42860-019-00041-0) Effect of heating on the XRD patterns. The hkl indices of the reflections belonging to the vermiculite structure are shown. The d values for the (00l) reflections before and after heating are also shown and the shifts are marked with arrows
In order to compare the new analyses with literature data, the structural formulae (Table 4) were calculated on the basis of 22 oxygens and assuming all iron to be ferric. The elemental changes in the crystal structure are represented in Fig. 8. Si, Al, and Fe increased progressively, whereas Mg decreased slightly during the vermiculitization at Sokli. These changes are largely similar to those described by Moon et al. (1994).

By using the calculation based on 22 oxygens for the structural formula (Table 4 and Fig. 8), the following trends can be seen: tetrahedral Al decreases with increasing octahedral Al (Fig. 8a), total Al increases with octahedral Al (Fig. 8b), Fe increases with increasing octahedral Al (Fig. 8c), Mg decreases with increasing octahedral Al (Fig. 8d), K decreases with increasing octahedral Al (Fig. 8f), and K decreases with increasing Ca (Fig. 8f).

DISCUSSION AND CONCLUSIONS

During the transformation from mica to vermiculite, numerous intermediate mixed-layer phases with various compositions and ordering are formed. As the vermiculite formation process is gradual, lamellae where potassium and ferrous ion concentrations are smaller than those of phlogopite can exist. Vermiculite differs from micas by not containing K+ and Fe(II), and having a significant amount of Fe(III) and water. The process of the alteration of mica to vermiculite is conceptualized in the literature as a replacement of K+ by Ca2+, Mg2+, and Al3+, oxidation of Fe(II) to Fe(III), and a loss of octahedral cations (Foster 1963; Rieder et al. 1998; Valášková et al. 2018).

An important general feature that illustrates the vermiculitization process is a decrease in K and an increase in water content (Table 4). Another feature is the increase of octahedral Al reported by Moon et al. (1994) and Toksoy-Köksal et al. (2001). The results comply with these characteristics and one can argue that the change from phlogopite to vermiculite occurs in the following way: due to weathering, K is released from the interlayer and Fe is converted from Fe(II) to Fe(III). Removal of K leaves a negative charge which is partly compensated by Ca-bearing fluids from the weathering carbonatite, which bring Ca2+ ions into the interlayer. The octahedral sheet is occupied initially by Fe(II), Mg, and Al, but as the vermiculitization proceeds, Mg is leached out and leaves the system. The negative charge left after loss of Mg is compensated partly by the conversion of Fe(II) to Fe(III). This compensation is insufficient and, consequently, Al is transferred from the tetrahedral sheet to the octahedral sheet and charge balance is partially reached. This sequence is similar to those presented in studies by Nemecz (1981), Banfield & Eggleton (1988), Moore & Reynolds (1989), and Moon et al. (1994).

Calculating the structural formula on the basis of 22 oxygens may not represent the exact distribution of cations in the structural layers, perhaps as a consequence of focusing on cations and ignoring the differing amounts of interlayer water content in the calculations. For example, the absolute SiO2 content in all analyses appears to be similar (Table 4). If Si is

| Natural | Heated | Natural | Heated |
|---------|--------|---------|--------|
| Sokli1  | 2.948  | 2.463   | 2.507  |
| Sokli7  | 2.946  | 2.464   | 2.509  |
| Sokli8  | 2.945  | 2.461   | 2.517  |

Figure 7. Back-scattered electron image of the mica sample Sokli10: (a) cross section and (b) surface. The lighter areas represent greater potassium contents in the mica.
fixed in the tetrahedral sites, Si should be constant, and Al should fill the rest of the available vacancies in the tetrahedra. The Al that does not fit the tetrahedral sites is situated in the octahedral sites. Fluctuation of the water content (and other fluids) and the form of water (H2O or OH−) may produce negatively charged vacancies in the lattice that can be occupied by cations that do not fit a theoretical filling of the structure based on 22 oxygens. The sum of octahedral cations does not exceed 6 in any of the points measured, which is likely due to the leached-out Mg.

This study adds information identifying vermiculite as a possible future commodity at the Sokli carbonatite complex and may improve the economic value of the Sokli intrusion by identifying the vermiculite at Sokli as a valuable resource. Mica clay from the regolith overlying the Sokli complex in northeastern Finland was characterized using EPMA, XRD, and TG-DSC techniques. Vermiculite is the dominant mica mineral in the deposit. The combined results indicate that the investigated phases correspond to vermiculite, with subordinate amounts of phlogopite. The presence of phlogopite is indicated by varying K2O contents (0.0–8.0 wt.%) and by XRD diffractograms showing a pattern typical of natural Mg-vermiculite. The TG and DSC results showed different thermal behavior in the samples investigated because of differences in chemical composition. The dehydration of Sokli vermiculite took place in two steps over the temperature range 25–170°C. The dehydration-induced diminution of interlayer distance typical of vermiculite was observed clearly in the XRD patterns measured before and after the heating of samples for 1 h at 600°C.

Table 4 EPMA results (wt.%) and structural formula calculated by the method of Deer et al. (1992)

|       | 1   | 2   | 3   | 4   | 5   | 6   | 7   | 8   | 9   | 10  | 11  | 12  | 13  | 14  |
|-------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| SiO2  | 41.91 | 41.08 | 41.42 | 41.72 | 40.42 | 41.94 | 42.97 | 41.07 | 40.48 | 42.33 | 41.69 | 41.76 | 41.01 | 41.39 |
| TiO2  | 0.65  | 0.56  | 0.64  | 0.57  | 0.77  | 0.53  | 0.51  | 0.58  | 0.61  | 0.65  | 0.56  | 0.47  | 0.52  | 0.55  |
| Al2O3 | 12.98 | 14.10 | 14.21 | 13.18 | 13.84 | 13.53 | 13.52 | 13.35 | 13.33 | 13.20 | 13.36 | 14.35 | 13.70 |
| Fe2O3 | 4.17  | 5.54  | 5.59  | 5.04  | 5.16  | 5.87  | 5.66  | 6.94  | 6.65  | 6.82  | 6.40  | 6.35  | 6.76  | 6.41  |
| MnO   | 0.03  | 0.02  | 0.03  | 0.02  | 0.06  | 0.04  | 0.02  | 0.05  | 0.04  | 0.08  | 0.04  | 0.03  | 0.05  | 0.05  |
| MgO   | 25.08 | 24.38 | 23.43 | 20.83 | 21.01 | 19.31 | 19.48 | 17.52 | 18.52 | 18.81 | 18.09 | 20.09 | 19.98 | 19.07 |
| CaO   | 0.02  | 0.03  | 0.22  | 0.90  | 0.45  | 1.40  | 1.43  | 1.30  | 2.18  | 1.23  | 1.29  | 1.88  | 1.16  | 1.42  |
| Na₂O  | 0.50  | 0.79  | 0.74  | 0.95  | 0.74  | 0.02  | 0.08  | 0.03  | 0.03  | 0.01  | 0.00  | 0.00  | 0.00  | 0.02  |
| K₂O   | 7.24  | 6.73  | 5.93  | 5.35  | 4.77  | 0.28  | 0.23  | 0.06  | 0.05  | 0.04  | 0.03  | 0.02  | 0.01  | 0.01  |
| Tot.  | 88.42 | 93.86 | 92.74 | 87.27 | 89.77 | 82.05 | 82.36 | 80.90 | 83.40 | 82.35 | 80.62 | 82.70 | 85.16 | 82.91 |

Number of cations based on 22 oxygen

Si 5.95 5.85 5.92 6.05 6.18 6.28 6.30 6.36 6.29 6.30 6.36 6.16 6.23 6.30

IVAl 2.05 2.15 2.08 1.95 1.82 1.72 1.70 1.64 1.71 1.70 1.64 1.84 1.77 1.70

SUM(IV)² 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.00

VAl 0.12 0.18 0.28 0.37 0.52 0.71 0.72 0.80 0.70 0.69 0.77 0.56 0.72 0.74

Ti 0.07 0.06 0.06 0.06 0.08 0.06 0.06 0.07 0.07 0.07 0.07 0.05 0.06 0.06

Fe³⁺ 0.45 0.58 0.59 0.57 0.56 0.67 0.65 0.81 0.75 0.78 0.75 0.73 0.75 0.73

Mn 0.00 0.00 0.00 0.00 0.01 0.01 0.01 0.00 0.01 0.01 0.00 0.01 0.01 0.01

Mg 5.31 5.10 4.93 4.65 4.50 4.40 4.41 4.04 4.16 4.27 4.18 4.56 4.38 4.29

SUM(VI)³ 5.95 5.92 5.87 5.65 5.68 5.85 5.84 5.72 5.69 5.83 5.78 5.91 5.91 5.82

Mg 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00

Ca 0.00 0.00 0.03 0.14 0.07 0.23 0.23 0.22 0.35 0.20 0.21 0.31 0.18 0.23

Na 0.14 0.21 0.20 0.28 0.21 0.01 0.02 0.01 0.01 0.00 0.00 0.00 0.00 0.01

K 1.31 1.20 1.07 1.02 0.87 0.05 0.05 0.01 0.01 0.01 0.00 0.01 0.00 0.00

SUM(i.lyr)⁴ 1.45 1.42 1.30 1.44 1.15 0.29 0.30 0.24 0.37 0.21 0.22 0.31 0.19 0.24

1 Numbers indicate the number of analyzed points
2 Tetrahedral sheet
3 Octahedral sheet
4 Interlayer

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Figure 8. Compositional variation of atoms per formula unit (atoms p.f.u.) diagrams for investigated mica from Sokli (black squares) compared with analyses from Moon et al. (2008) (red dots): (a) IVAl vs. VIAl; (b) Al(tot) vs. VIAl; (c) Fe(tot) vs. VIAl; (d) Mg vs. Mg(tot); (e) K vs. VIAl; and (f) K vs. Ca. In the graphics, IVAl is tetrahedral aluminum, VIAl is octahedral aluminum, Al(tot) is total aluminum, and VI Mg is octahedral magnesium.
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