# OXYGEN ISOTOPE CHANGES DURING MICA ALTERATION

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Abstract – Oxygen isotope analyses ( $\delta O^{18}$ ) of micas that were artificially depleted in K<sup>+</sup> indicate little or no isotope exchange during the transformation. The oxidation of iron in K-depleted, iron-rich micas by H<sub>2</sub>O<sub>2</sub> treatment resulted in 1.6 to 4.6% decrease in  $\delta O^{18}$  due to the fact that the equilibrium fractionation factor is less than the initial difference between the starting  $\delta O^{18}$  of the fluid and micas. The oxygen isotope ratio of a saponite formed by the weathering of phlogopite showed a 9.7% increase in  $\delta O^{18}$  due to authigenic recrystallization. These results suggest that oxygen isotope ratios can be used to determine the nature of chemical transformation during the weathering of mica to vermiculite and/or smectite.

Key Words-Mica, Oxygen isotopes, Potassium, Recrystallization, Saponite, Weathering.

# INTRODUCTION

Oxygen and hydrogen isotope compositions of ocean sediments (Savin and Epstein, 1970) and Quaternary soils (Lawrence and Taylor, 1971) that consist largely of authigenic minerals are in isotopic equilibrium with the coexisting waters. According to Savin and Epstein (1970) detrital clays, however, show no major oxygen isotope exchange with the coexisting waters, but these authors were unable to make a definitive statement about hydrogen isotope exchange of detrital clays in sea water. Unweathered minerals in soil profiles developed on igneous and sedimentary rocks also undergo no appreciable oxygen or hydrogen isotope exchange with meteoric water in the weathering environment (Lawrence and Taylor, 1972). Thus, O18/O16 ratios may be used to distinguish authigenic minerals from detrital components because detrital minerals originating from igneous and metamorphic rocks have lower O18/O16 ratios than most minerals formed at low temperatures (Hoefs, 1980). The above statement, however, may not be applicable to assemblages found at high latitudes and high altitudes where meteoric waters are depleted in O<sup>18</sup> (S. M. Savin, Case-Western Reserve University, Cleveland, Ohio, personal communication, 1984).

Quaternary soils commonly contain appreciable quantities of detrital micas and their transformation products, as well as varying amounts of authigenic minerals. Detrital micas transform to vermiculite and then to minerals of the smectite group (Jackson *et al.*, 1952). These mineralogical changes result in the replacement of K<sup>+</sup> from micas by hydrated cations, with concurrent decrease in layer charge. The decrease in layer charge results from compositional changes such as the oxidation of iron, the ejection of structural cations such as Mg<sup>2+</sup>, Fe<sup>3+</sup>, and Al<sup>3+</sup> (Veith and Jackson, 1974), and the incorporation of Si in the layers (Sridhar and Jackson, 1974). The clays formed from the alteration of micas swell in water (Sridhar et al., 1972) and commonly have aluminous and/or ferruginous coatings (Roth et al., 1969). Although the changes in chemical composition during the above transformations have been studied by many workers, the extent of isotope exchange between the aluminosilicate framework and the associated waters has received little or no study. The objective of the present study is to determine the extent of oxygen isotope exchange in various structural layers during the weathering of micas to expanding layer silicates, both in nature and in the laboratory.

# **EXPERIMENTAL**

# Materials

Phlogopite and saponite were separated from the Hills Pond peridotite, Woodson County, Kansas. Here, phlogopite in the parent peridotite has weathered in part to saponite through a vermiculite intermediate (Jackson and Sridhar, 1974). Phlogopite of igneous or metamorphic origin from near Perth, Lanark County, Ontario, was obtained as large sheets from Ward's Nat-

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ural Science Establishment, Rochester, New York. Biotite from Bancroft, Ontario, Canada ( $<50 \ \mu m$  size, courtesy, A. D. Scott), biotite from an unknown locality (20–250  $\mu$ m), and an iron-rich mica, lepidomelane (20– 250  $\mu$ m) were also obtained as large sheets from Ward's Natural Science Establishment. Muscovite mica (5– 10  $\mu$ m) from Norway was also used.

# Sample preparation

Phlogopite flakes (>500  $\mu$ m) were isolated from the Kansas peridotite by wet sieving and hand picking and then washed with water to remove the adhering fine particles. The saponite ( $<0.2 \mu m$ ) was separated from the peridotite by dispersing the sample in water and centrifuging the dispersion. The clay-size fraction was treated with pH 5 N NaOAc, H<sub>2</sub>O<sub>2</sub>, and citrate-bicarbonate-dithionate to remove calcite, organic matter, and free iron oxides, respectively (Jackson, 1956), as described below. The purity of this material was tested by X-ray powder diffraction (XRD) and quantitative mineralogical analysis (Jackson and Sridhar, 1974). Samples of phlogopite (Ontario), biotite (Ward's), lepidomelane, and muscovite (Norway) were wet ground in a blender and the respective size fractions obtained by sedimentation and centrifugation (Jackson, 1956). Biotite (Bancroft) was ground dry and screened into different size fractions.

Deferration. Saponite and the iron-rich biotite micas were treated to remove iron oxide and oxyhydroxide coatings (i.e., deferration) using sodium dithionate  $(Na_2S_2O_4)$  as a reducing agent, sodium bicarbonate as a buffer, and sodium citrate as a chelating or complexing agent for ferrous and ferric iron (Jackson, 1956). After the treatment, the samples were washed free of all salts.

NaOAc and  $H_2O_2$  treatments. To eliminate carbonates and organic matter in the <0.2- $\mu$ m fraction of saponite, the sample was washed with 1 N NaOAc at pH 5 and digested in 30%  $H_2O_2$ . The Kansas phlogopite (>500  $\mu$ m) and the various mica samples were not treated this way because carbonates and organic matter were not apparent in these samples under the light microscope.

K-depletion of micas. To simulate natural vermiculitization, the various micas were treated with sodium tetraphenylboron (NaTPB) in 1 N NaCl solution (Scott and Smith, 1966). This process lowers the K<sup>+</sup> activity in the suspending liquid by the precipitation of potassium tetraphenyl boron. Muscovite mica was heated to 550°C prior to the NaTPB treatment to enhance the K-depletion process (Scott *et al.*, 1972). Heating muscovite at 550°C should not affect its isotope exchange properties because this temperature is well below its dehydroxylation temperature (Jackson, 1956). The NaTPB treatment results in a Na<sup>+</sup>-saturated vermiculite with  $H_2O$  in the interlayers. The NaTPB treatment was carried out to determine the extent of isotope exchange with water which entered the interlayers. The artificial K<sup>+</sup>-depletion process also resulted in the partial oxidation of iron in iron-rich micas (Sridhar and Jackson, 1974).

 $Fe^{2+}$ -oxidation treatment. The iron-rich micas, biotites and lepidomelane, after K<sup>+</sup>-depletion, were digested with 30% H<sub>2</sub>O<sub>2</sub> at ~90°C for 6 hr to oxidize most of the structural Fe<sup>2+</sup> present (Jackson, 1956). The exact amount of iron oxidized by the H<sub>2</sub>O<sub>2</sub> treatment, however, was not determined.

Oxygen isotope determination. Oxygen from the micas and their weathering products was extracted by reacting 10 to 20 mg of solid with BrF<sub>5</sub> at 550°C in nickel reaction vessels for 12–14 hr (Clayton and Mayeda, 1963). The evolved O<sub>2</sub> was then converted to CO<sub>2</sub> for mass spectrometric analysis by combustion with graphite (Taylor and Epstein, 1962). Analyses of CO<sub>2</sub> for oxygen isotopes were made using an isotope-ratio mass spectrometer. The isotopic data are reported as  $\delta O^{18}$  values in per mil (‰), where  $\delta O^{18}$  is defined as:

$$\delta O^{18} = \left[ \frac{O^{18}/O^{16} \text{ sample}}{O^{18}/O^{16} \text{ standard}} - 1 \right] \times 1000,$$

with Standard Mean Ocean Water (SMOW) (Craig, 1961) as a reference.

# **RESULTS AND DISCUSSION**

The oxygen isotope ratios of various micas and their transformation products are presented in Table 1. Simple exchange of interlayer K<sup>+</sup> by Na<sup>+</sup> in the artificially weathered phlogopite and muscovite caused no significant change in the oxygen isotope composition. In two iron-rich micas (Bancroft biotite and Ward's lepidomelane), the  $\delta O^{18}$  value decreased significantly after K<sup>+</sup>-depletion. This decrease in iron-rich micas may have been related to the partial iron oxidation that took place (Table 1) during K<sup>+</sup>-depletion as found by Sridhar and Jackson (1974). The effect of iron oxidation on the oxygen isotope ( $\delta O^{18}$ ) ratio was investigated further by treating the samples with  $H_2O_2$  (Table 1). Oxidation of structural iron should have increased with the  $H_2O_2$  treatment, but the exact amount was not determined. The H<sub>2</sub>O<sub>2</sub> treatment resulted in a large  $\delta O^{18}$  decrease in iron-rich micas. This decrease in  $\delta O^{18}$ accompanying  $Fe^{2+}$  oxidation by  $H_2O_2$  treatment may reflect oxygen isotopic exchange between the clay and the interlayer water which was introduced during K<sup>+</sup>depletion; however, there is no clear cut correlation between the Fe<sup>2+</sup> contents of these micas (as reported by Sridhar and Jackson, 1974) and the decrease in  $\delta O^{18}$ by Fe2+ oxidation. The distilled water used in the K-depletion experiments had a  $\delta O^{18}$  value of -8 to -12%. The oxidation of iron results in the ejection of Fe<sup>3+</sup> and/or Mg<sup>2+</sup> cations from octahedral sites in order to balance the charge (Veith and Jackson, 1974). Deuteration experiments have suggested that the hydroxyls become exchangeable even at room temperature (unpublished results). The extent of exchange of hydroxyls with interlayer water is not known. A substantial hydroxyl exchange with the low  $O^{18}$  water might have led to a lower  $\delta O^{18}$  value after oxidation.

A second explanation for the lower  $\delta O^{18}$  value after K-depletion is related to the hydrolysis of iron ejected from the octahedral sites. The ejected iron could have hydrolyzed with the interlayer water, resulting in low  $\delta O^{18}$  phases in the interlayer positions and hence an overall lower  $\delta O^{18}$  value for the depleted mica. In an equilibrium system, this decrease in  $\delta O^{18}$  for the mica transformation product results only if the equilibrium fractionation factor is less than the initial difference between the starting  $\delta O^{18}$  of the fluid and solid. The fact that the iron-rich micas were originally rich in O18 suggests some low-temperature interaction prior to treatment. Because the starting micas were already enriched in O<sup>18</sup>, some depletion took place during the K+-replacement. If these micas were originally depleted in O<sup>18</sup> (as expected for an igneous or metamorphic source), significant enrichment could have resulted. The effect of  $H_2O_2$  treatment on the  $\delta O^{18}$  of K-depleted, iron-poor phlogopite and muscovite needs to be investigated before the effect of iron oxidation on  $\delta O^{18}$ can be confirmed.

The lack of oxygen isotope exchange during the simple mineral transformation involving Na<sup>+</sup> and water exchange with K<sup>+</sup> for the iron-poor micas, phlogopite, and muscovite suggests that the oxygen isotope ratio can be an indicator of partial authigenic recrystallization, i.e., breakdown of framework metal-oxygen bonds leading to the removal and incorporation of cations in transformation products. Authigenic recrystallization by the removal of tetrahedral Fe and Al and the incorporation of Si has been proposed (Sridhar and Jackson, 1974) as the mechanism for the decrease in layer charge in the natural transformation of phlogopite to saponite. If the layer-charge decrease during this natural transformation was indeed caused by the removal and incorporation of structural ions with the breakdown of framework metal-oxygen bonds, this transformation might have led to oxygen isotope exchange. To test this possibility, the oxygen isotope ratios of phlogopite and its natural weathering product, saponite, were determined (Table 1). Phlogopite from Kansas has a  $\delta O^{18}$  value of 6.7‰ which indicates its high-temperature (igneous) origin. On the other hand, the saponite transformation product of this phlogopite has a significantly higher  $\delta O^{18}$  value of 16.4‰. This drastic increase (9.7‰) of  $\delta O^{18}$  in saponite compared to the parent phlogopite indicates extensive oxygen isotope exchange during weathering. Because this phlogopite contains little iron and because no deuteric alteration appears to have taken place, the change in 
 Table 1. Oxygen isotope abundance of micas and their transformation products.

	Isotopic	Fe <sup>2+</sup>
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Sample	SMOW	100 g
Micas and artificially weathered micas		
Phlogopite, Ontario	16.7	NA
Phlogopite, Ontario, K <sup>+</sup> -depleted	17.2	NA <sup>1</sup>
Muscovite, Norway	11.3	NA
Muscovite, Norway, K <sup>+</sup> -depleted		
after heating to 550°C	11.4 <sup>2</sup>	NA <sup>1</sup>
Biotite, Bancroft, Ontario <sup>3</sup>	11.1 <sup>2</sup>	NA <sup>1</sup>
Biotite, Bancroft, Ontario,		
K <sup>+</sup> -depleted	10.2	31
Biotite, Bancroft, Ontario,		
K <sup>+</sup> -depleted and Fe oxidized		
with $H_2O_2$	9.5	$ND^{i}$
Biotite, Ward's Natural Science		
Establishment <sup>3</sup>	12.0	NA
Biotite, Ward's Natural Science		
Establishment, K <sup>+</sup> -depleted	11.8	17
Biotite, Ward's Natural Science		
Establishment, K <sup>+</sup> -depleted		
and Fe oxidized with $H_2O_2$	7.7	ND
Lepidomelane, Ward's Natural		
Science Establishment <sup>3</sup>	11.5	$NA^1$
Lepidomelane, Ward's Natural		
Science Establishment	10.1	56
Lepidomelane, Ward's Natural		
Science Establishment, <sup>3</sup>		
K <sup>+</sup> -depleted and Fe oxidized		
with $H_2O_2$	6.9	ND
Phlogopite and its natural weathering p	roduct	
Phlogopite, Woodson County,		
Kansas	6.7 <sup>2</sup>	NA
Saponite formed from phlogopite,		
Woodson County, Kansas <sup>4</sup>	16.4 <sup>2</sup>	NA <sup>1</sup>

 $^{1}$  NA = not applicable; ND = not determined; the values for Fe<sup>2+</sup> oxidation were taken from Sridhar and Jackson (1974).

<sup>2</sup> Average of duplicates.

<sup>3</sup> Treated with citrate-bicarbonate-dithionate.

 $^4$  Treated with  $H_2O_2,\,pH$  5 N NaOAc and citrate-bicarbonate-dithionate.

 $\delta O^{18}$  from phlogopite to saponite could be due to oxygen isotope exchange resulting from the breakdown of framework metal-oxygen bonds. Thus, these oxygen isotope data support the Sridhar and Jackson's (1974) ' ypothesis of authigenic recrystallization by the re-1..oval of tetrahedral Fe<sup>3+</sup> (and Al<sup>3+</sup>) and incorporation of Si<sup>4+</sup> during the transformation of phlogopite to saponite. The chemical process of phlogopite transformation to saponite with the accompanying oxygen isotopic change can be summarized as follows:

# Phlogopite + Si<sup>4+</sup>

# $\underbrace{\overset{Oxygen \text{ isotope exchange}}{\longleftarrow} Saponite + Fe^{3+}(Al^{3+}) + K^+$

The removal of  $Fe^{3+}$  or  $Al^{3+}$  and the incorporation of Si in the tetrahedral sheets of this mica led to a breakdown in framework metal-oxygen bonds, and this breakdown facilitated oxygen isotopic exchange.

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# CONCLUSIONS

Little or no oxygen isotope exchange occurred in vermiculite by artificial K<sup>+</sup>-depletion. The oxidation of iron in Fe-rich micas by  $H_2O_2$  treatment led to a decrease in the  $\delta O^{18}$  value probably due to oxygen isotope exchange with the water introduced into the interlayers and/or by forming low-O<sup>18</sup> oxide phases by the hydrolysis of ejected iron in the interlayers. This depletion of  $\delta O^{18}$  in mica transformation products was likely due to the fact that the equilibrium fractionation factor was less than the initial difference between the starting  $\delta O^{18}$  of the fluid and mica. Authigenic recrystallization during the mineral transformation of phlogopite to saponite was supported by an increase in the  $\delta O^{18}$  ratio.

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# REFERENCES

- Clayton, R. N. and Mayeda, T. K. (1963) The use of bromine pentafluoride in the extraction of oxygen from oxides and silicates for isotopic analysis: *Geochim. Cosmochim. Acta* 27, 43-52.
- Craig, H. (1961) Standard for reporting concentrations of deuterium and oxygen 18 in natural waters: Science 133, 1833-1834.
- Hoefs, J. (1980) Stable Isotope Geochemistry: 2nd ed., Springer-Verlag, Berlin, 208 pp.
- Jackson, M. L. (1956) Soil Chemical Analysis-Advanced Course: 5th printing, 1969, published by the author, Department of Soil Science, University of Wisconsin, Madison, Wisconsin, 894 pp.

Jackson, M. L., Hseung, Y., Corey, R. B., Evans, E. J., and Vanden Heuvel, R. C. (1952) Weathering sequence of clay-size minerals in soils and sediments: II. Chemical weathering of layer silicates: *Soil Sci. Soc. Amer. Proc.* 16, 3-6.

- Jackson, M. L. and Sridhar, K. (1974) Scanning electron microscopic and X-ray diffraction study of natural weathering of phlogopite through vermiculite to saponite: *Soil Sci. Soc. Amer. Proc.* **38**, 843–847.
- Lawrence, J. R. and Taylor, M. P., Jr. (1971) Deuterium and oxygen-18 correlation: clay minerals and hydroxides in Quaternary soils compared to meteoric waters: *Geochim. Cosmochim. Acta* 35, 993–1003.
- Lawrence, J. R. and Taylor, M. P., Jr. (1972) Hydrogen and oxygen isotope systematics in weathering profiles: *Geochim. Cosmochim. Acta* 36, 1377–1393.
- Roth, C. B., Jackson, M. L., and Syers, J. K. (1969) Deferration effect on structural ferrous-ferric iron ratio and CEC of vermiculites and soils: *Clays & Clay Minerals* 17, 253– 264.
- Savin, S. M. and Epstein, S. (1970) The oxygen and hydrogen isotope geochemistry of ocean sediments and shales: *Geochim. Cosmochim. Acta* 34, 43-63.
- Scott, A. D., Ismail, F. T., and Locatis, R. R. (1972) Changes in interlayer potassium exchangeability induced by heating micas: in *Proc. Int. Clay Conf., Madrid, 1972, 2, J. M.* Serratosa, ed., Division De Ciencias C.S.I.C., Madrid, 467– 479.
- Scott, A. D. and Smith, S. J. (1966) Susceptibility of interlayer potassium in micas to exchange with sodium: in *Clays* and *Clay Minerals, Proc. 14th Natl. Conf., Berkeley, California, 1966, S. W. Bailey, ed., Pergamon Press, New York,* 69-81.
- Sridhar, K. and Jackson, M. L. (1974) Layer charge decrease by tetrahedral cation removal and silicon incorporation during natural weathering of phlogopite to saponite: *Soil Sci. Soc. Amer. Proc.* 38, 847–850.
- Sridhar, K., Jackson, M. L., and Syers, J. K. (1972) Cation and layer charge effects on blister-like osmotic swelling of micaceous vermiculite: *Amer. Mineral.* 57, 1832–1848.
- Taylor, H. P., Jr. and Epstein, S. (1962) The relationship between O<sup>18</sup>/O<sup>16</sup> ratios in coexisting minerals of igneous and metamorphic rocks. *Geol. Soc. Amer. Bull.* 73, 461– 480.
- Veith, J. A. and Jackson, M. L. (1974) Iron oxidation and reduction effects on structural hydroxyl and layer charge in aqueous suspension of micaceous vermiculites: *Clays & Clay Minerals* 22, 345–353.

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Резюме — Изотопный анализ кислорода ( $\delta O^{18}$ ) в образцах слюды, содержание K<sup>+</sup> в которых было искуственно уменьшено, указывают на незначительный или нулевой изотопный обмен во время превращения. Результатом окисления железа в K-обедненных, обогащенных железом слюдах, при помощи обработки перекисью было уменьшение  $\delta O^{18}$  на 1,6 до 4,6‰. Это было вызвано тем, что величина равновесного коэффициента разделения была меньше, чем разница начальных значений  $\delta O^{18}$  для течи и слюд. Отношение изотопов кислорода для сапонита, образованного путем выветривания флогопита показало 9,7‰ увеличение значения  $\delta O^{18}$ , вызванное аутогенной рекристаллизацией. Эти результаты указывают на то, что отношения изотопов кислорода могут быть использованы для вермикулита и/или смектита. [E.G.]

**Resümee**—Sauerstoffisotopen-Analysen ( $\delta O^{18}$ ) von Glimmern, denen K<sup>+</sup> künstlich entzogen wurde, zeigen geringen oder gar keinen Isotopenaustausch während der Umwandlung. Die Oxidation von Eisen in K<sup>+</sup>-verarmten, Eisen-reichen Glimmern durch H<sub>2</sub>O<sub>2</sub>-Behandlung führte zu einer Abnahme von 1,6 bis 4,6‰ bei  $\delta O^{18}$ , aufgrund der Tatsache, daß der Gleichgewichtsfraktionierungsfaktor kleiner ist als der ursprüngliche Unterschied zwischen dem Ausgangswert  $\delta O^{18}$  der Flüssigkeit und dem der Glimmer. Das Sauerstoffisotopenverhältnis eines Saponits, der durch Verwitterung von Phlogopit gebildet wurde, zeigte eine Zunahme von 9,7‰ des  $\delta O^{18}$ -Wertes aufgrund autigener Rekristallisation. Diese Ergebnisse deuten darauf hin, daß die Sauerstoffisotopenverhältnisse dazu verwendet werden können, die Art der chemischen Umwandlung während der Verwitterung von Glimmer zu Vermiculit und/oder Smektit zu bestimmen. [U.W.]

**Résumé** – Des analyses d'isotope oxygène ( $\delta O^{18}$ ) de micas dont on a artificiellement retiré K<sup>+</sup> ont indiqué peu ou pas d'échange d'isotopes pendant la transformation. L'oxidation du fer dans des micas riches en fer, privés de K par traitement H<sub>2</sub>O a resulté en une diminution d'1,6 à 4,6‰ de  $\delta O^{18}$  à cause du fait que le facteur de fractionation d'équilibre est plus petit que la différence initiale entre le  $\delta O^{18}$  de départ du fluide et des micas. La proportion d'isotope oxygène d'une saponite formée par l'altération d'une phlogopite a montré une augmentation de 9,7‰ de  $\delta O^{18}$  à cause de la recristallisation authigénique. Ces résultats suggèrent que les proportions d'isotope oxygène peuvent être utilisées pour déterminer la nature de la transformation chimique pendant l'altération du mica en vermiculite et/ou en smectite. [D.J.]