Atomic-resolution transmission electron microscope evidence for the mechanism by which chlorite weathers to 1:1 semi-regular chlorite-vermiculite

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ABSTRACT

Atomic-resolution transmission electron microscope (TEM) images reveal that IIbb ($\beta = 97^\circ$) Mg,Al,Fe-chlorite from Koongarra, Australia, transforms to vermiculite via a range of intermediate chemical and structural states. Semi-quantitative analysis of contrast in atomic-resolution images of layer silicates with $\sim 1.4$ nm basal spacings indicate that the interlayers range from brucite-like to having $\sim 0.3$–$0.6$ interlayer cations per formula unit. Octahedral cations (predominantly Mg and Fe) tend to be removed from every second interlayer, leading to semi-regular 1:1 interstratifications of chlorite-vermiculite. Further loss of interlayer cations is accompanied by partial to complete interlayer collapse in the vacuum of the TEM. Resulting intergrowths of chlorite and semi-regular 1:1 chlorite-vermiculite retain the primary chlorite orientation, morphology, and sense of octahedral tilt in 2:1 layers. Although vermiculitization is a continuous process that occurs by a solid-state mechanism, the reaction involves important structural modifications. Atomic-resolution [010] images indicate initial loss of interlayer cations is accompanied by $\sim a/3$ shifts of 2:1 layers and cations in brucite-like interlayers. Displacements of interlayer cations change the interlayer stacking from IIbb to Iab and shift of the following 2:1 layer converts it from lab to laa. Displacements are driven by the lower energy of $a$-type interactions when vacancies occur in sites above tetrahedral cations. Shift of a 2:1 layer alters the subsequent interlayer from IIbb to IIab. Stabilization of every slightly altered second interlayer by introduction of $a$-type stacking explains development of semi-regular 1:1 chlorite-vermiculite interstratifications. Displacements occur before significant modification of interlayer electron density can be detected in high-resolution images. This observation is consistent with previously reported inhibition of layer shifts by low interlayer charge. Layer displacement may occur by an elastic process (no rupture of bonds within the 2:1 layer) at the tip of the growing vermiculite portion of the intergrowth. Removal of cations from the chlorite-vermiculite junction may be facilitated by rapid diffusion along the vacancy-rich interlayer. Mg is removed in solution, Fe is precipitated locally in aggregates of nanocrystalline Al-, Si-, and P-bearing goethite.

INTRODUCTION

Geochemical and biogeochemical cycles near the Earth’s surface are greatly impacted by weathering reactions that release cations to solution and produce clay minerals. Most mineral weathering studies involve measurement of dissolution rates under conditions that preclude formation of clay minerals. In contrast, some natural weathering reactions produce abundant clays that develop topotactically on primary mineral surfaces (e.g., Eggleton 1984; Banfield and Barker 1994; Hochella and Banfield 1995). Because of structural coherence between the primary and secondary phases, polymers may be inherited by weathering products. The mechanisms of solid state or partially solid-state transformations are different from dissolution-reprecipitation and thus require different kinetic models.

The best evidence that low-temperature reactions can involve considerable structural inheritance comes from studies of naturally weathered layer silicates. Lattice-fringe TEM images indicate a layer-by-layer biotite-to-vermiculite transformation mechanism (Banfield and Eggleton 1988; Kogure and Murakami 1996) accompanied by gradual reduction in the K content of the interlayer (Kogure and Murakami 1996). Direct formation of vermiculite from biotite can also be inferred from experimental studies showing incongruent dissolution (e.g., Acker and Bricker 1992; Kalinsowski and Schweda 1996), which may be especially important at pH > 4.0 (Acker and Bricker 1992). This paper examines the detailed structural rearrangements that accompany the weathering of chlorite to vermiculite. From these data, we propose a reaction mechanism and explain the formation of semi-regular interstratified chlorite-vermiculite.

SAMPLES AND EXPERIMENTAL METHODS

Samples examined in this study are from diamond drill core from a weathered portion of the Koongarra uranium
deposit, within the Alligator River Uranium Field in Northern Australia. Geological details were summarized by Snelling (1992). Murakami et al. (1996) provide a cross section of the deposit. The silicate mineralogy of the Koongarra deposit was described by Komninou and Sverjenski (1995). Hydrothermal conversion of muscovite-2M to a K-poor, Si-rich muscovite-2M, and formation of chlorite from solution and by replacement of altered muscovite was proposed. The quartz chlorite schist examined in this study was subjected to weathering for more than one Ma (Airey et al. 1986). Murakami et al. (1996) characterized variably weathered samples by X-ray diffraction and electron microprobe analysis (EMPA). Weathering products of Fe-rich (chamosite) and Mg-rich clinohlore) chlorite identified by X-ray diffraction (XRD) included chlorite-vermiculite intergrade (as defined by Wilson 1987), interstratified chlorite-vermiculite, vermiculite, and kaolinite.

Samples examined in this study are DDH 3-160 (37.4 m depth), DDH 3-106 (24.8 m depth), DDH 3-105 (24.5 m depth), DDH 3-103 (24.1 m depth), and DDH 4-101 (23.6 m depth). Samples range from very slightly to moderately weathered.

Approximately 30 μm thick slices were removed from petrographic thin sections and thinned to electron transparency by argon ion milling. Specimens were coated lightly with carbon and examined in a JEOL 2010 transmission electron microscope with a point resolution of 0.2 nm. Analytical electron microscope (AEM) analyses were collected with a Li-drifted Si detector and processed with a JEOL analyzer. Images were recorded on Mitsu-bitui film at magnifications of 150 000x to 800 000x using exposure times of ~1–6 s.

Selected areas of TEM negatives were digitized and processed to remove noise and the image of amorphous material by rotational filtering (NCEM package of Kilaas and Siddnei implemented within Digital Micrograph version 2.5, Gatan Ltd.). Details in filtered and unfiltered images were interpreted to provide atomic information by comparison with simulated images calculated using MacTempus software (Total Resolution Co.).

RESULTS

Hydrothermal mineralogy

Scherzer defocus atomic-resolution images down [100] (Fig. 1, top) clinohlore show spotty contrast associated with both octahedral sheets (small arrows in Fig. 1, top). Image simulations from chlorite with completely triocatahedral character have a continuous dark stripe over the octahedral sheets whereas images from chlorite with diocatahedral sheets have spotty contrast. Thus, contrast in Figure 1, top, indicates that both octahedral sheets in this chlorite have some diocatahedral character. AEM analyses indicate that this is mostly due to significant Al (also see EMPA analyses in Murakami et al. 1996).

The 02l and 04l rows in most [100] selected area electron diffraction patterns (SAED) patterns from unwethered chlorites show maxima superimposed upon continuous streaks. However, in some regions, the stacking is regular (Fig. 1, top). The 20l reflections in SAED patterns are sharp. The abundant polytype in all samples is the IIb polytype (β ~97°), although the Ibb polytype (β = 90°) is present also. Energy-dispersive X-ray (EDX) analyses show that chlorite Mg:Fe ratios are variable. A continuum of compositions exists between clinohlore and chamosite.

HRTEM images reveal that most chlorite crystals contain occasional serpentine layers (Fig. 1, bottom). These
typically occur as single layers, occasionally in semi-periodic intergrowths with chlorite. Small areas of dozyite (Bailey et al. 1995) were identified, and longer period arrangements (e.g., chlorite, chlorite, serpentine, as in Fig. 1, bottom) occur locally. Rarely, packets containing up to about 10 serpentine layers were detected.

Large crystals of 2M$_1$ mica occur in some areas (Fig. 2). The two-layer periodicity is apparent in SAED patterns (Fig. 2b). K contents are impossible to quantify by AEM due to loss of K during analysis. This phase is characterized by high Si:Al ratios compared with ideal muscovite and 1–5% Mg and Fe. Careful examination of a series of high-resolution images recorded from a several micrometer-wide area, including the analysis region showed no chlorite or serpentine layers. Probably, Mg and Fe are incorporated in the muscovite structure.

Image simulations (Fig. 3) show that the darkest spots in Figure 2 correspond to projected pairs of tetrahedra, the dioctahedral sheet of the 2:1 layer is resolved as slightly weaker spots midway between the tetrahedra. The regular two-layer alternation of the stacking vector within the 2:1 layer (parallel and at 60° to the beam) is apparent in Figure 2. The contrast associated with interlayer K is relatively weak compared with contrast expected from ideal muscovite (see simulations in Fig. 3). Experimental images closely match calculated images at K contents of 0.3 and 0.5 per formula unit (pfu). The match between the simulation for 0.4 K pfu and experimental image is shown in Figure 2d.

**Weathering of chlorite**

Chlorite that is structurally and compositionally different from typical unweathered material is characterized by streaks along the 20l and 40l rows in SAED patterns (Fig. 4). Although streaking along 02l and 04l rows in layer silicates is well understood (e.g., Bailey 1988a), streaking of h0l is uncommon and has not been discussed in detail previously (e.g., Bailey 1988a, 1988b).

Figure 5a shows an atomic-resolution Scherzer defocus image recorded down [010] from a region characterized by pronounced streaking along 20l. Contrast was manipulated in Figure 5b to emphasize discrete spots. Details in Figure 5b show that the $\beta$ angle is not consistent over
Figure 4. The [010] selected-area electron diffraction patterns from (a) fresh, and (b), (c) increasingly altered chlorite. Note the streaks along 20l in (c).

more than a few unit cells. Arrows between contrast maxima reveal displacements both of interlayer cations and 2:1 layers. The magnitude of the smallest displacement measured from the images is \(-0.2 \text{ a} (\sim 0.106 \text{ nm})\).

Typically chlorite structurally modified by layer displacements is characterized by highly variable contrast associated with the brucite-like interlayer in Scherzer defocus images (e.g., Figs. 5b, 6, 7, and 8). In some areas, reduction of contrast of brucite-like interlayers correlates with layer displacement along \(a^*\) (best illustrated in Fig. 7) and superposition of tetrahedral sites across low-contrast interlayers (Fig. 7b).

Lower contrast interlayers tend to develop adjacent to darker contrast interlayers and in some areas this leads to a semi-regular alternation (see Fig. 7). Note that displacements (arrows in Fig. 7) correlate with lower contrast interlayers. In some interlayers, octahedral sites (projected M3 and M4 sites) are clearly resolved as spots (Fig. 8). Image simulations for clinochlore (Fig. 9) indicate that contrast changes due to interlayer occupancy are not apparent until the M3 and M4 sites are about half occupied. Distinct spots occur when the interlayer site occupancy approaches 0.2 (0.6 cations pfu; however, simulations with interlayer occupancies of 0.1 are not substantially different). On the basis of visual comparison with experimental images, the interlayer in partly weathered chlorite (that retains the 1.4 nm basal spacing in the vacuum) may range from brucite-like (as in ideal chlorite) to near vermiculite-like.

Interstratifications with interlayers that are more brucite-like alternate in a semi-periodic fashion with layers characterized by 1.0 nm basal spacings (note the obvious \(\sim 2.4 \text{ nm} \) separation of higher contrast interlayers in Fig. 10). The spacings between these interlayers is commonly variable. In Figure 10, for example, more brucite-like interlayers are separated by \(\sim 2.37 \text{ to } 2.63 \text{ nm}\). The largest spacings are observed where low-contrast relics remain in the more vermiculite-like interlayer (see arrow in Fig. 10).
EDX analyses of regions consisting of mixtures of 1.4 and 1.0 nm phases show similar Al:Si ratios but lower Mg and Fe contents than chlorite. XRD analyses of clays from DDH 3-105 indicated exchange behavior consistent with a vermiculite-like phase but thermal behavior resembling chlorite (Murakami et al. 1996). This behavior is consistent with layer charge intermediate between chlorite and vermiculite. Units with modified contrast but 1.4 nm spacings are interpreted as chlorite-vermiculite intergrade and those with \(\sim 1.0\) nm basal spacings (Figs. 10, 11, and 12) are probably vermiculite.

Figure 11 is Scherzer defocus [010] zone axis image showing interlayers with more brucite-like contrast (site occupancies probably \(> 0.5\)) every fourth interlayer. On the left and right sides, the two interlayers bracketed by brucite-like interlayers have highly modified contrast. These layers probably have occupancies approaching, but not equal to, those of vermiculite (see Fig. 9). In the center of Figure 11, the brucite-like interlayers enclose layers with \(\sim 1.0\) nm basal spacings. The interlayer occupancy of this material is too low to prevent collapse in the TEM vacuum. Thus, the layer spacings in Figures 10 and 11 are artifacts of differential collapse, reflecting subtle differences in interlayer occupancies. The sequence in Figure 11 approaches chlorite-vermiculite-vermiculite (but is periodic only over the area shown).

In DDH 3-103 and DDH 4-10, layer silicates are a fine-scale intergrowth of chlorite, semi-regular 1:1 chlorite-vermiculite interstratifications, and minor vermiculite (Fig. 12, top). SAED patterns show broad (00\(l\)) reflections with diffuse maxima corresponding to 1.2, 0.8, and 0.6-0.7 nm (Fig. 12, bottom). These regions preserve the (often bent) crystal shapes of primary chlorite as well as the primary grain orientations, which in some cases involve intergrowths of packets of chlorite with [010] parallel to [100].

Despite complications due to the variable Fe:Mg ratio of primary chlorite, AEM analyses suggest alteration is associated with significant loss of Fe and Mg, resulting in aluminous vermiculite. Packets of 1.0 nm material co-exist with interstratified chlorite and chlorite-vermiculite in some regions of more altered samples. These occasionally contain uncollapsed layers with more brucite-like character (Fig. 13). The \(\beta\) angle for the 1.0 nm phase is
Figure 9. Simulations of high-resolution images of chlorite (11bb structure of Rule and Bailey 1987) calculated using parameters for a JEOL 2010 HRTEM with Cs = 0.5 mm at close to Scherzer defocus. The left column shows results for a sample 5 nm thick and the right column for a sample 7 nm thick. The defocuses for the top, middle, and bottom rows in each group are $-40$, $-42$, and $-44$ nm (a) M3 and M4 sites fully occupied, (b) each M3 and M4 site is half occupied (1.5 interlayer cations pfu), (c) each M3 and M4 site 20% occupied (0.6 interlayer cations pfu).

Figure 10. Rotationally filtered Scherzer defocus [100] image of interstratified chlorite-like and vermiculite-like material. Note the three chlorite-vermiculite units (~2.4 nm separation of more brucite-like interlayers). The arrow indicates a vermiculite-like interlayer that contains a small area of low contrast material that is associated with increase in the separation of enclosing more brucite-like interlayers from ~2.37 to 2.63 nm.

Figure 11. Rotationally filtered Scherzer defocus [010] image of interstratified chlorite-like and vermiculite-like layers superimposed over the unfiltered image. Arrows indicate the positions of interlayers associated with layers with 1.4 nm basal spacings. Such sequences do not persist over large distances.

close to 97° (Fig. 13 inset; but 90° in some cases). The polytype for the vermiculite-like phase was not uniquely determined, but SAED data are consistent with Iaa, which is the common structure identified in XRD studies of vermiculite (de la Calle and Suquet 1987).

Iron precipitation results in local concentrations of semi-oriented aggregates of goethite nanocrystals between packets of altered layer silicates. These have an approximately topotactic orientation ([001] goethite parallel to [010] chlorite-vermiculite and $b^*$ goethite parallel to $a^*$ chlorite-vermiculite), implying Fe-dominated octahedral strips in goethite are parallel to octahedral sheets
**Figure 12.** (top) Scherzer defocus [100] image of typical interstratified chlorite-vermiculite (CV) and chlorite-like (C) material in the most weathered samples examined. Low contrast, wedge-shaped areas (e.g., ?) may be space produced by collapse across vermiculite-like interlayers in the vacuum. (bottom) SAED pattern for an area of material including that in top figure. Note that fuzzy spots along 00l at 1.22 and 0.8 nm are expected diffraction maxima associated with mixed layer chlorite-dioctahedral vermiculite (bottom).

**Figure 13.** Rotationally filtered Scherzer defocus [010] image of the 1.0 nm phase containing isolated interlayers with some brucite-like character. (Inset in upper right corner) Fourier transform of (a) showing discrete 20l reflections and β ~ 97°.

in associated layer silicate. Topotactic packets of kaolinite are also present in some regions.

**Discussion**

**Mineralogy of the least-altered samples**

The mineralogy of these samples is similar to that reported by Komninou and Sverjensky (1995). The Si-rich muscovite-2M₁ was shown to be K-deficient relative to ideal muscovite (~0.4 K per formula unit). This composition is similar to that proposed by Komninou and Sverjensky (1995). Probably this illite-like phase was formed by a continuous process that did not require dis-

solution and reprecipitation of the basic structure. Illitization may have occurred either during weathering or a low-temperature hydrothermal event.

**Weathering**

Structural and chemical data are interpreted to indicate a continuous range of site occupancies in the brucite-like interlayer of weathered chlorite. In addition, the sense of shift in the 2:1 layer is inherited by the vermiculite-like product. Thus, we propose that chlorite was vermiculitized by a continuous, solid-state mechanism. The lower activation energy for direct structural modification compared with bulk dissolution and recrystallization can be explained based on the high degree of structural correspondence between chlorite and vermiculite (the 2:1 layer is almost identical).

The result that natural chlorite weathering is incongruent (at least under some conditions) differs from conclu-
sions reached by many previous workers. Nagy (1995) noted that although most laboratory studies report nonstoichiometric dissolution early in experiments, dissolution is congruent in the later stages. The more recent results of Malmstrom et al. (1996) for chlorite dissolution at near-neutral pH are similar to those reviewed by Nagy (1995).

In contrast to dissolution studies where all constituents are released into solution, the fate of constituents initially present in the chlorite examined here is highly variable. Si and Al are retained in the intergrown chlorite- and vermiculite-like material. A significant fraction of the Mg and Fe are released, Mg to solution and Fe to form secondary Fe-oxides. These results for chlorite are analogous to those for naturally weathered biotite (Banfield and Eggleton 1988; Kogure and Murakami 1996) and from some experimental dissolution studies (Acker and Bricker 1992). The complexity evident in some kinetic studies of layer silicate dissolution (see discussion in Kalinowski and Schweda 1996) is understandable in view of these data.

Despite the continuous nature of compositional changes, results show that all structural characteristics of chlorite are not inherited. An early step in the vermiculitization reaction involves shifts of some 2:1 layers with respect to adjacent layers. Where these shifts occur frequently but nonperiodically, there is no fixed value of \( \beta \), resulting in streaking (or spottiness) along 20l.

Direct measurements from atomic-resolution images show that the smallest shift between 2:1 layers is \(-0.2 \ a\), equivalent to a shift of \(-0.3 \ a\) in the opposite direction. The vector 0.3a is close to the a/3 vector that separates a from b positions in the interlayer. Figure 14 shows the consequences of a series of a/3 displacements of interlayer cations and a 2:1 layer. The gray line connects tetrahedra (gray circles) in the lower 2:1 layer, and the solid line connects tetrahedra in the 2:1 layer above the interlayer octahedral tilt so that the chlorite unit, including the interlayer region (i.e., II\(ab\)) chlorite. If the interlayer cations (circles with +) are displaced downward by a/3 (arrows), M3 sites superimpose directly onto tetrahedral sites in the 2:1 layer below (Fig. 14a). Note that this displacement alters the interlayer octahedral tilt so that the chlorite unit, consisting of this interlayer and the underlying 2:1 layer, is now lab (interlayer hydroxyls remain fixed).

If the 2:1 layer above (gray circles linked by solid black lines) relocates by an upward shift of a/3 (Fig. 14b), the structure in this region becomes Iaa, as expected for vermiculite. Note that pseudohexagonal cavities oppose each other, so the upper tetrahedral cations project directly onto the interlayer cations and the tetrahedral cations below (as in Fig. 7b).

A predominant product of chlorite weathering in the samples described here is a regular 1:1 interstratification of chlorite and vermiculite. Johnson (1964) also identified this as the product of chlorite weathering and suggested it formed from a two-layer chlorite polytype by selective alteration of one of the two interlayers. However, no evidence exists for a two-layer polytype in chlorite studied here (or in similar weathered chlorite examined by Churchman 1980).

Formation of 1:1 chlorite-vermiculite must involve changes in stability of every second brucite-like interlayer as a consequence of modification of the first. Figure 14c shows that the \(-a/3\) displacement of brucite-like layers observed in [010] images (transforms the first interlayer to laa) also changes the subsequent interlayer, converting it from IIab to IIaa. These displacements are summarized in Figure 15 (similar displacements are required to create aa character in the interlayer of IIbb chlorite).

The apparent stability of the alternate interlayers (IIab) suggests these have lower interlayer site occupancies than original IIbb brucite-like interlayers. However TEM observations of semi-regular 1:1 interstratifications of vermiculite and more chlorite-like layers suggest that further cation loss from these more brucite-like interlayers is arrested. This may result from the introduction of a-type character, which reduces the energy of the partly vermiculitized interlayer region (i.e., IIab \( \rightarrow \) Iaa lab), and thus reduces the driving force for further displacements. Consequently, interlayer shift is the vehicle by which modification of one layer affects the reactivity of the next. In conclusion, semi-regular chlorite-vermiculite formation is partly a consequence of the kinetic controls, which favor direct structural transformation over dissolution-reprecipitation. It is also partly the result of an energy minima resulting from increased stability of alternate interlayers due to reduction in repulsion between interlayer cations and cations in tetrahedral sites.

Shifting entire 2:1 layers during a weathering reaction
are perhaps surprising, given that the justification for solid state transformation is minimization of structural rearrangement. However, analogous layer shifts take place during (diagnostic) vermiculite cation exchange reactions. For example, de la Calle and Suquet (1987) note that cation exchange in vermiculite is accompanied by changes in the relative positions of the 2:1 layers. Although these cation exchange reactions differ in detail, such observations show that 2:1 layer displacements can take place in response to changes in the interlayer easily and at low temperature.

Similar layer shifts accompany the solid-state serpentine-to-chlorite reaction in metamorphic rocks (Banfield and Bailey 1996). Presumably these shifts are driven by the greater stability of type \(b\) vs. type \(a\) interactions between octahedral interlayer cations and cations in the tetrahedral and octahedral sites in the 2:1 layer of trioctahedral chlorite. Banfield and Bailey (1996) suggested that the shift of the 2:1 layer occurs sequentially as the new 2:1 layer forms, with the displacement accommodated at the junction by elongation of tetrahedral-O bonds. In this case, and in the vermiculitization of chlorite, changes in the relative positions of 2:1 layers occur elastically. We predict that other solid-state layer silicate alteration reactions (e.g., muscovite or biotite-1M ↔ chlorite-IIbb) should involve analogous displacements (and polytypic changes) to those documented here.

The relative stability of \(a\)-type interactions in the \(1aa\) structure of vermiculite was attributed to vacancies in energetically unfavorable sites directly above tetrahedral cations (Bailey 1988b). Conversely, high site occupancies in the brucite-like interlayer of chlorite favor \(b\)-type stacking with sites that avoid superposition over tetrahedral cations. As chlorite vermiculitization increases the number of vacancies on interlayer sites, the stability of \(b\)- vs. \(a\)-type stacking is reversed. In the chlorite-to-vermiculite reaction, these cationic interactions are clearly sufficiently important to trigger displacement of interlayer cations and the associated 2:1 layers.

Cation exchange studies of vermiculite show that layer displacements are inhibited where interlayer charge is low (de la Calle and Suquet 1988). This is attributed to reduction in cationic repulsions originating between the 2:1 layer and interlayer. This is consistent with the findings of this study. These findings show that layer displacement occurs early in the chlorite-to-vermiculite weathering reaction when interlayer occupancies are still close to (but lower than) those of chlorite.

Our results show that Fe was transported many micrometers from reaction sites. We do not know the pH or redox state of solutions at grain boundaries and in narrow fluid conduits at the time of reaction. However, the solutions were probably chemically and physically distinct from bulk meteoric H2O. Presumably, Fe was moved from reaction sites inside crystals by interlayer diffusion, transported by solution as Fe\(^{2+}\) to more oxidized regions, and precipitated. This resulted in formation of aggregates of nanocrystals of goethite, which nucleated and grew topotactically on layer silicate surfaces.

Although most studies concur that vermiculitization is associated with loss of Mg and Fe, the role of Fe oxidation is unclear (Ross and Kodama 1976; Proust et al. 1986). If iron is removed from chlorite as Fe\(^{2+}\), as suggested above, charge imbalance due to oxidation does not trigger loss of interlayer cations and other changes associated with vermiculitization.

We cannot unequivocally establish the role of Fe oxidation in promoting vermiculitization. However, it is useful to examine what other charge balance changes could promote loss of interlayer cations. A fundamental difference between chlorite and vermiculite is the presence of H2O molecules in place of hydroxyl groups surrounding interlayer cations. Our data imply a range of interlayer compositions exist involving any mixture of M\(_{\text{c}}\cdot\text{(OH)}\(_x\) and M\(_{\text{c}}\cdot\text{n H}_2\text{O}\) (where \(x\) and \(y\) indicate dioctahedral character in the brucite-like and vermiculite interlayer, respectively, and \(n\) is the number of H2O molecules). Thus, an increase in \(a_{\text{H}}\), may result in sequential protonation of hydroxyls in the chlorite interlayer, with charge balance maintained by loss of interlayer cations, not unlike ex-
changes proposed to operate in other mineral weathering reactions (e.g., Na\(^+\) ↔ H\(^+\) and Ca\(^+\) ↔ 2H\(^+\) in weathering of feldspars). A strong pH dependence of alteration rates [analogous to that reported by Kalinowski and Schweda (1996) for trioctahedral micas] would support this suggestion.

Formation of vermiculite, kaolinite, and goethite

Some samples contain highly altered regions of vermiculite, kaolinite, and goethite. Loss of Mg and Fe from octahedral sites in chlorite results in vermiculite layers with considerable dioctahedral character. Although the original distribution of Al between octahedral sites in the 2:1 layer and brucite-like interlayer is not known, [100] zone axis images suggest sites in both octahedral sheets have limited dioctahedral character. If the resulting vermiculite has Mg in the interlayer, as suggested by Murakami et al. (1996), exchange of Al and Mg between the 2:1 layer and interlayer may occur. Chlorite-vermiculite may be converted to vermiculite by removal of cations from remaining brucite-like interlayers (with some structural rearrangement). Alternatively, vermiculite formation from chlorite-vermiculite may involve dissolution-reprecipitation, as is probably the case for formation of kaolinite.

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