Cs-exchange in birnessite: Reaction mechanisms inferred from time-resolved X-ray diffraction and transmission electron microscopy

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ABSTRACT

We have explored the exchange of Cs for interlayer Na in birnessite using several techniques, including transmission electron microscopy (TEM) and time-resolved synchrotron X-ray diffraction (XRD). Our goal was to test which of two possible exchange mechanisms is operative during the reaction: (1) diffusion of cations in and out of the interlayer or (2) dissolution of Na-birnessite and reprecipitation of Cs-birnessite. The appearance of distinct XRD peaks for Na- and Cs-rich phases in partially exchanged samples offered support for a simple diffusion model, but it was inconsistent with the compositional and crystallographic homogeneity of (Na,Cs)-birnessite platelets from core to rim as ascertained by TEM. Time-resolved XRD revealed systematic changes in the structure of the emergent Cs-rich birnessite phase during exchange, in conflict with a dissolution and reprecipitation model. Instead, we propose that exchange occurred by sequential delamination of Mn oxide octahedral sheets. Exfoliation of a given interlayer region allowed for wholesale replacement of Na by Cs and was rapidly followed by reassembly. This model accounts for the rapidity of metal exchange in birnessite, the co-existence of distinct Na- and Cs-birnessite phases during the process of exchange, and the uniformly mixed Na- and Cs-compositions ascertained from point analyses by selected area electron diffraction and energy dispersive spectroscopy of partially exchanged grains.

Keywords: Cation exchange, cesium, birnessite, synchrotron, X-ray diffraction, transmission electron microscopy

INTRODUCTION

Groundwater contamination by radionuclides (including U, Pu, Np, and Cs) is a serious problem at several national laboratories that were involved in the production of components for nuclear weapons in the United States, such as the Hanford Site in Washington state (McKinley et al. 2001). Leaks from high-level waste (HLW) storage tanks in the 200 Area of the Hanford Site have released appreciable quantities of $^{137}$Cs into the vadose zone, and migration of this contaminant has extended to depths that are significantly greater than expected (Serne et al. 2001a, 2001b). Radioactive $^{137}$Cs is a fission product of irradiated U and Pu with a half-life of 35.7 years (Zachara et al. 2002). Because it is highly soluble, $^{137}$Cs can be extremely mobile in soil environments (Bostick et al. 2002). However, its rate of migration through soils is difficult to model because the transport of $^{137}$Cs depends on numerous factors, particularly fluid composition and soil type (Almgren and Isaksson 2006). For example, several researchers have demonstrated that $^{137}$Cs will readily sorb to various aluminosilicate clay minerals (Comans et al. 1991; Sutton and Sposito 2001; Bostick et al. 2002; Zachara et al. 2002), dramatically inhibiting transport through the subsurface.

In the present study, we explored the interaction of dissolved Cs ions with layered Mn oxides, which are ubiquitous in a wide range of soils, from arid desert varnishes to temperate soil precipitates (Waychunas 1991; Post 1992, 1999; Yang et al. 2003), including the Ringold Formation, which underlies the Hanford formation and comprises a mixture of poorly consolidated clays, silts, sands, and gravels (Fredrickson et al. 2004). Even when Mn oxides occur at the 1 wt% level or lower in soils, these phases can act as the controlling players in contaminant migration. The high reactivity of these minerals can be attributed to several factors. Many Mn oxide phases occur as particles that are only a micrometer in diameter or smaller, particularly when they grow authigenically within soils. Consequently, the ratio of reactive surface area to volume is extremely high (Murray 1974, 1975). In addition, Mn oxides can occur in various structural topologies (Fritsch et al. 1997), and many of the phases that are commonly found in soils have structural architectures (e.g., layer-type or tunnel-type) that are especially amenable to solid-state diffusion (Balachandran et al. 2002).

Birnessite-like phases are the most common natural phyllosilicates. The birnessite structure (Fig. 1) consists of sheets of edge-sharing Mn$^{2+}$O$_6$ octahedra where Mn$^{2+}$ or vacancies substitute for Mn$^{4+}$ in the octahedral layers, resulting in a net negative layer charge, which is balanced by various hydrated cations in the interlayer region (commonly Na and Ca). Many studies have quantitatively demonstrated that Mn oxides (particularly birnessite) are sinks for a host of transition metals (Loganathan and Burau 1973; Singh and Subramanian 1984; Burns et al. 1985; Nicholson and Eley 1997), and even transuranic radionuclides have been shown to exhibit a strong affinity for Mn oxides (Triay et al. 1991; Duff et al. 2001, 2002; Powell et al. 2006).
In addition to sorption on the surfaces of Mn oxides, it long has been known that dissolved cations will exchange for structural cations within the interlayers and tunnels of the Mn oxide phases (Golden et al. 1986a; Tsuji et al. 1992; Kuma et al. 1994). The exchange process bears strongly on the efficiency of birnessite for the inhibition of contaminant migration because different pathways for cation exchange may offer variable capacities for Cs sequestration in birnessite.

Here we consider three reaction mechanisms that might govern the exchange of Cs for Na in birnessite: diffusion, dissolution/re-precipitation, and delamination.

In the diffusion model, the substitution of one cation for another occurs by solid-state transport of one species into the interlayer region as the other species diffuses out. This scenario is the traditional paradigm for understanding cation exchange, and it implies that the Mn-O octahedral sheets are passive scaffolds and undergo no structural change during the diadochic reaction. In this model, the exchange is constrained by the counter-ions being redistributed by diffusion until equilibrium is achieved (Helfferich 1962).

In contrast, Putnis (2002) argues that many mineral replacement reactions take place primarily by dissolution/re-precipitation processes. For example, O’Neil and Taylor (1967) used $^{18}$O to monitor the reaction between albite (NaAlSi$_3$O$_8$) and KCl in hydrothermal solutions to produce sanidine (KAlSi$_3$O$_8$), and they found that during the reaction the oxygen isotope distribution re-equilibrates in the product. They attributed this result to the breaking of Si-O and Al-O bonds. Similarly, O’Neil (1977) demonstrated by stable isotope experiments that simple cation exchange reactions take place primarily by dissolution/re-precipitation, and delamination.

For delamination, an exfoliation, layer (Liu et al. 2000). No researchers have yet reported evidence for delamination of Na-birnessite upon washing. An exfoliation, or delamination/reassembly (or restacking) process, is involved in the synthesis of various important materials, including nanocomposites (Yang et al. 2004). If the cation exchanges in our experiments occur through delamination, then presumably the Mn-O octahedral layers (and thus the Mn-O bonds) remain intact but exfoliate and then re-order, or self-assemble.

To evaluate these three possible pathways for Cs exchange, we monitored crystallographic changes in the birnessite structure in real time using time-resolved synchrotron X-ray diffraction coupled with Rietveld structure analysis. We also examined the partial and complete run products of Cs-exchanged Na-birnessite using a variety of techniques, including X-ray diffraction (XRD), inductively coupled plasma atomic emission spectroscopy (ICP-AES), and transmission electron microscopy (TEM).

**Experimental Methods**

**Na-birnessite synthesis**

Synthetic birnessite was prepared using the methods outlined in Lopano et al. (2007), based on protocols in Golden et al. (1986b). A chilled (~5 °C) 250 mL solution of 5.5 M NaOH was reacted with 200 mL of a chilled 0.5 M MnCl$_2$ solution at room temperature while oxygen was bubbled through the resulting Mn(OH)$_2$ suspension at a rate of over 1.5 L/min. Similar to the procedure outlined in Kuma et al. (1994), the precipitate was filtered through a 0.1 µm Millipore filter, washed multiple times, and stored in water until aliquots were removed and air-dried at room temperature for use in the synchrotron X-ray diffraction experiments and laboratory batch experiments.

**Synchrotron X-ray diffraction**

Approximately 1 mg samples of Na-birnessite were loaded into 0.7 mm quartz capillaries and held in place by cotton or glass wool on either side of the samples. Individual capillaries were inserted into a flow-through apparatus similar to the SECReT's cell (Lee et al. 1998, 2000; Parise et al. 2000). In successive experiments, solutions of 0.001, 0.01, and 0.05 M CsCl at pH 7 were passed through the sample cell at an average rate of ~1 drop/min, and X-ray diffraction patterns were collected every 2 min with a MAR345 full imaging plate detector for periods of 1 to 3 h depending on concentration. The detector-sample distance was roughly 175 mm for all experiments, which allows us to achieve a typical 2θ range of ~4–50°. A sample of the stacked diffraction patterns for the 0.01 M Cs-exchange is shown in Figure 2.

The reacted samples from the synchrotron experiments were pressed flat on a carbon stub and analyzed using a Cameca SX-50 electron probe microanalyzer (EPMA) to determine whether any Na remained in the exchanged solid. No Na was detected in most of the Cs-exchanged samples. However, a trace amount of Na remained in the 0.001 M Cs-exchanged sample. The small sample size and the fine-grained, powdered nature of the material prevented accurate quantitative chemical analysis by electron beam methods, but the absence of detectable Na indicated that exchange was complete for all reactions except for the 0.001 M exchange.

Rietveld refinements (Rietveld 1969) of the birnessite structures were performed using the EXPGUI interface (Toby 2001) of the general structure analysis.
The batch synthesized Cs,Na-birnessite sample was prepared using a method modified from Kuma et al. (1994), by reacting 1 g of Na-birnessite with MnCl₂ solution in 50 mL centrifuge tubes. These centrifuge tubes were placed on a shaker table and reacted for 24 h. The resulting product was then centrifuged 5 min in time. Patterns were collected at NSLS with a wavelength of 0.9218 Å.

The initial structure parameters for the unreacted Na-birnessite came from the triclinic structure described in Post et al. (2002). The background intensities for the X-ray diffraction patterns were fit using up to 26 terms of a linear interpolation function. The peak profiles were modeled by a pseudo-Voigt profile function as parameterized by Thompson et al. (1987), with asymmetry corrections by Finger et al. (1994) and microstrain anisotropic broadening terms of Stephens (1999).

During initial cycles of refinement only the background, scale, peak profile, and unit-cell parameters were allowed to vary. The position of the O atom in the Mn-O sheet was then refined. Difference-electron Fourier (DELF) maps were calculated to determine the combined O (water) and Cs interlayer atom positions. For the Cs-exchange experiments described in this study, it was found that two different split interlayer sites were required to describe the interlayer electron density, where these sites could contain Cs and/or O. Following refinements of the interlayer atom positions, occupancy factors and isotropic atomic displacement parameters were allowed to vary. The position of the O atom in the Mn-O sheet, and unit-cell parameters were refined. Following refinements of bond distances is displayed in Table 3.

The final χ² values for end-member Cs-birnessite ranged from 0.7 to 0.9 (Table 1). Because amorphous components (glass capillary and water) contributed very high background intensities to our patterns, the calculated estimated standard deviations (e.s.d.) for our patterns were unreasonably large. Consequently, the weighting factors were unreasonably low, generating values for χ² that consistently fell below unity. Nevertheless, the overall excellence of the goodness-of-fit parameters attests to the accuracy of the refined structure models. Refinement results for the end-member exchanged birnessites at the different concentrations are presented in Table 1; atom positions are included in Table 2; and a summary of bond distances is displayed in Table 3.

### Batch exchange experiments

Laboratory batch experiments were performed to synthesize an incompletely exchanged Cs,Na-birnessite and a completely exchanged Cs-birnessite. The batch exchange method was modified from that specified in Golden et al. (1986a, 1986b). For the completely exchanged Cs-birnessite, 25 mg of Na-birnessite were reacted with 40 mL of a 1 M CsCl solution in 50 mL centrifuge tubes. These centrifuge tubes were placed on a shaker table and reacted for 24 h. After 24 h, the exchange product was centrifuged and 40 mL of new solution was added. This process was repeated over a period of 5 days for a total of 200 mL of solution used in exchange. The reacted sample was then rinsed with distilled deionized water with subsequent centrifugation five times to remove excess salt. The sample was filtered with 0.1 μm Millipore filters and allowed to air-dry before being ground lightly under alcohol with a mortar and pestle.

The incompletely exchanged Cs,Na-birnessite sample was prepared using a method modified from Kuma et al. (1994), by reacting 1 g of Na-birnessite with 40 mL of 0.5 M CsCl solution in 50 mL centrifuge tubes. These centrifuge tubes were placed on a shaker table and reacted for 24 h. The resulting product was rinsed with distilled deionized water, with subsequent centrifugation five times to remove the excess salt. The solution then was filtered and air-dried before being lightly ground under alcohol.

The batch synthesized Cs-birnessites were analyzed using a variety of chemical analyses and microscopy techniques to ensure that a completely exchanged end-member was achieved, and to quantify the concentrations of Na and Cs in the incompletely exchanged samples. The Na-birnessite starting material, the end-member Cs-birnessite, and the Cs,Na-birnessite were analyzed using a Leeman Labs PS3000UV inductively coupled plasma atomic emission spectrophotograph (ICP-AES). The samples were dissolved in reverse aqua regia, then heated to dryness and dissolved in HCl. The Mn and Na quantities were determined by ICP-AES, and the amount of Cs was determined by flame emission after spiking with 1 g of LiBO₂ per 100 mL. The results of these analyses reveal that the starting Na-birnessite has a chemical formula of Na₆₋₇(Mn₄/₅Mn₁/₅)O₄·1.5H₂O and confirmed the EDS and EPMA results indicating that the Na-birnessite was completely exchanged. The incompletely exchanged Cs,Na-birnessite contained half Na and half Cs, with the chemical formula: Na₆₋₇Cs₃₋₄(Mn₄/₅Mn₁/₅)O₄·1.5H₂O. The chemical formulae were derived for each
Synchrotron X-ray diffraction
   
   Lattice variations with Cs-exchange. As Cs exchanged for Na in birnessite, the change in the basal spacing was sufficiently large that distinct peaks for Cs- and Na-birnessite became resolvable. Consequently, we refined 2 phases—Na-rich and Cs-rich birnessite—for each diffraction pattern. Our refinement of the Na-rich birnessite (until the fraction of the Na-rich phase decreased to a point at which the structure could no longer be refined) revealed that the lattice parameters remained constant within errors throughout the exchange process.

   Unlike the Na-rich phase, Cs-rich birnessite structurally evolved over the course of the exchange. The variations in the unit-cell parameters of the Cs-rich phases are plotted as a function of time in Figures 3, 4, and 5 for each ionic strength of the CsCl solution. The increase in unit-cell volume with progressive exchange correlates with an expansion along c; the a and β parameters, on the other hand, decreased. As the reaction progressed, small diffraction peaks appeared near the 002 reflection. These are likely superstructure reflections and were excluded from the refinements. The final end-member Cs-birnessite diffraction patterns (Fig. 6) showed significant decreases in the ratio of 001:002 peak intensities relative to that of Na-birnessite. Our simulations of powder X-ray diffraction patterns for different birnessite compositions suggested that this behavior is consistent with the exchange of heavy X-ray scatterers into the birnessite interlayer.

   Configuration of Cs-rich interlayer. In each Rietveld refinement of the Cs-rich member, the interlayer Cs cations and water molecules were modeled using a split O site, as suggested by the

   TEM of batch samples
   
   Because synthetic birnessite crystallites typically measured several micrometers or smaller in diameter and only 250 to 600 Å in thickness (Post and Veblen 1990), TEM was employed to study individual crystals. An EDS-capable JEOL 2010 TEM with a LaB₆ source was operated at an accelerating voltage of 200 keV and beam current typically of ~111 nA. The birnessite samples were dispersed in alcohol (typically methanol) and sonicated in an attempt to break up agglomerated fine particles. The birnessite particles were supported on 3.0 mm, 300 mesh holey carbon-coated copper grids. Samples were mounted in a Gatan double tilt holder. Selected area electron diffraction (SAED) patterns and EDS analyses were collected along traverses from the edges to the centers of individual grains of the incompletely exchanged Cs,Na-birnessite sample to determine the compositional homogeneity of each grain.

   RESULTS

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electron density calculated in the DELF maps (Fig. 7). Oxygen was chosen as proxy for H₂O, Cs, and Na species due to the partial occupancies of the disordered interlayer atom positions. It is not unusual for birnessite phases to exhibit positional disorder in the interlayer (Post et al. 2002; Lopano et al. 2007). Johnson and Post (2006) found that several metal-exchanged birnessite specimens contain two to three structurally different water sites, presumably related to disordering of cations in the interlayer. The waters associated with the relatively large Cs⁺ atom likely have more distorted geometries to coordinate the larger cation into the interlayer plane (Johnson and Post 2006). Because Cs⁺ has low

**Figure 4.** Changes in unit-cell parameters over time for the 0.01 M Cs-exchange reaction with Na-birnessite. The $b$, $\alpha$, and $\gamma$ parameters remained constant within error with exchange. The error bars calculated in GSAS are smaller than the plotting symbols.

**Figure 5.** Changes in unit-cell parameters over time for the 0.05 M Cs-exchange reaction with Na-birnessite. The $b$, $\alpha$, and $\gamma$ parameters remained constant within error with exchange. The error bars calculated in GSAS are smaller than the plotting symbols.

**Figure 6.** Observed (crosses), calculated (solid line), and difference (line below) plots for the end-member X-ray diffraction patterns from ~4.5 to 49.3 °2θ for: (a) 0.001 M exchange Cs-birnessite; (b) 0.01 M exchange Cs-birnessite; (c) 0.05 M exchange Cs-birnessite All were refined in a triclinic unit cell (CT). Peak positions are indicated by vertical dashes.
hydration energy, it does not bind the water molecules as tightly as would a smaller, more highly charged cation.

Bostick et al. (2002) studied Cs-adsorption on various clay minerals using EXAFS and found that bond distances varied with the degree of disorder in Cs\(^{+}\) coordination environments. They found that when outer-sphere complexes form (i.e., when the cation retains its solvation shell), Cs-O bonds are consistent with those of Cs\(^{+}\) ions in aqueous solution. Thus, the bonds are short (on the order of 3.0–3.2 Å) because hydrated Cs\(^{+}\) ions typically exhibit a lower coordination than other Cs\(^{+}\) ions (Bostick et al. 2002). Partial dehydration of the Cs\(^{+}\) ion can result in inner-sphere complexes where a Cs-surface bond forms, and these Cs-O bond distances are longer (due to a higher coordination number) and can range from ~3.3 to ~4.8 Å (Bostick et al. 2002).

In our refinements of Cs-substituted birnessite, the two interlayer sites were separated by only ~1 Å; therefore, both sites cannot be occupied within the same unit cell. Most of the O\(_{oc}-O_{as}\) and O\(_{oc}-O_{at}\) distances (2.8–3.2 Å) (Table 3) are close to the values expected for Cs as an outer-sphere complex (remembering that 2/3 of the electron density at the site arises from Cs, so that O\(_{as}\) is modeling more Cs than H\(_2\)O). Some of the longer distances (on the order of 3.4 Å) might represent Cs atoms that are partially dehydrated. Cs\(^{VI}\)-O\(^{IV}\) bond distances calculated from Shannon (1976) are 3.1 Å, and therefore, interatomic distances that are much smaller than this value (e.g., ~2.6 Å in Table 3), likely represent hydrogen bonds between water molecules. In light of the degree of disorder in the Cs-birnessite interlayer, it is not possible to determine the spatial distribution of Cs\(^{+}\) cations and H\(_2\)O molecules as clearly as was achieved for interlayer K\(^+\) and Ba\(^{2+}\) in Lopano et al. (2007), but one could propose several possible ordering schemes based on occupancy factors and relative amounts of Cs and water.

**Kinetics of Cs-exchange.** The transformation from Na-birnessite to Cs-birnessite was more rapid in solutions with higher ionic strength, as seen in the shift of the 002 peak (Figs. 8 and 9) and the increase in the 001:002 peak height ratios during the exchange. The formation of end-member Cs-birnessite during exchange with the 0.05 M CsCl solution began as soon as the solution touched the sample, as is also apparent from the unit-cell parameter changes with time (Figs. 3, 4, and 5). Precipitation of Cs-birnessite in the presence of the 0.01 M CsCl occurred more slowly, but it commenced within the first 10 min of the experiment. For reactions involving a 0.001 M CsCl solution, the appearance of the Cs-birnessite peak required ~20 min to develop. These observations are consistent with those in Lopano et al. (2007) for K- and Ba-exchange.

**TEM analysis of batch samples**

**Compositional homogeneity.** TEM analysis of the birnessite samples was challenged by several factors. Particles were
consistently agglomerated in our suspension mounts, making it difficult to isolate individual grains. Moreover, variations in grain thickness also were evident (Fig. 10). Nevertheless, for batches of end-member Na- and Cs-birnessite as well as birnessite that had been partially exchanged, we identified single birnessite grains with optimal qualities for TEM investigation. For each grain, we collected SAED patterns and EDS analyses along traverses from the edges to the centers to determine the compositional homogeneity of the grain.

EDS spectra from numerous single grains of end-member Na- and Cs-birnessite confirmed that they were chemically homogeneous and pure. Significantly, the EDS spectra collected from the partially exchanged Cs,Na-birnessite samples revealed no compositional variations as a function of distance from the crystal edges, nor did the intermediate Cs,Na-birnessite compositions vary from grain to grain (Fig. 11). Thus, we found no evidence for chemical zonation, or reaction rims, as we might expect if diffusion was the operational exchange process. Surprisingly, we also discerned no evidence for the co-existence of compositionally distinct phases by EDS, as was shown to occur by XRD patterns of the same samples.

**Electron diffraction results.** Representative SAED patterns for the Na- and Cs-birnessite end-members and for a 50% substituted phase are shown in Figure 12. The presence of streaking is indicative of the structural disorder in each of the samples. Electron diffraction patterns of Na-birnessite exhibited streaks along the <110> directions, whereas streaking in Cs-birnessite was strongest along the <310> directions. In addition, SAED patterns of Cs-birnessite exhibited distinct superstructure reflections at 1/3 <310>. The d-spacing associated with this superstructure (4.33 Å) matches within experimental error the most intense of the satellite reflections revealed by synchrotron XRD (a broad peak from 4.0 to 4.3 Å). These observations are consistent with findings reported in Post and Veblen (1990), Manceau et al. (1992), Kuma et al. (1994), and Drits et al. (1998); although these authors differ in their speculations for the origin of the superstructures.

Interestingly, a comparison of the SAED patterns from our three sample compositions (Fig. 12) revealed that grains of the intermediate Cs,Na-birnessite exhibited features consistent with both Na-birnessite and Cs-birnessite. The Cs,Na-birnessite SAED pattern retained the streaking in the <110> direction (as in Na-birnessite), but it also included superstructure reflections at 1/3 <310> (as in Cs-birnessite). Although these superstructure
as a straightforward diadochic exchange in which birnessite platelets swapped hydrophilic Na cations out of the interlayer for dissolved K and Ba cations. If simple diffusion were the operative mechanism for Cs-exchange, one might hypothesize that partially exchanged birnessite platelets would exhibit a marked compositional heterogeneity because the rims of the platelets would serve as the active sites for exchange. Consequently, the platelet edges would be enriched in Cs, whereas the unreacted cores of the platelets might retain their interlayer Na until the reaction was complete.

Our real-time X-ray diffraction analyses of Cs-exchange in Na-birnessite would seem to support this model. Synchrotron XRD of the partially reacted birnessite provided definitive evidence for the co-existence of two birnessite phases with different interplanar 00l spacings, and thus, different Na and Cs concentrations. Surprisingly, however, our EDS and SAED analyses of partially substituted Na,Cs-birnessite were equally definitive in demonstrating the compositional homogeneity of individual platelets along a core-to-rim traverse. In the absence of distinct reaction rims for partially exchanged samples, we conclude that simple diffusion of dissolved Cs from the solution into the birnessite interlayer regions cannot account for the reaction textures that we observed.

**Discussion**

**Implications of XRD and TEM results for exchange mechanisms**

When K and Ba were substituted for Na in birnessite during the experiments of Lopano et al. (2007), only one phase could be resolved, and we assumed that the exchange occurred as an evolving single-phase solid solution. It is possible, however, that the substitution of K and Ba for Na into the birnessite interlayer actually generated biphasic mixtures, but the diffraction resolution of those experiments, even with synchrotron X-ray radiation, was inadequate to distinguish a true solid solution from co-existing end-members with similar structures. In contrast, when we exchanged Cs for Na in birnessite in the present investigation, the expansion of the unit cell was sufficiently large that two phases could be distinguished. We argue that the unambiguous appearance of co-existing Na-rich and Cs-rich birnessite phases provides a key for determining which of our proposed cation exchange mechanisms is correct.

**Cation exchange by diffusion.** In Lopano et al. (2007), we presumed that the exchange of K and Ba for Na in birnessite occurred through a simple diffusion mechanism. Because we did not resolve co-existing phases, we modeled the reaction as a straightforward diadochic exchange in which birnessite platelets swapped hydrophilic Na cations out of the interlayer for dissolved K and Ba cations. If simple diffusion were the operative mechanism for Cs-exchange, one might hypothesize that partially exchanged birnessite platelets would exhibit a marked compositional heterogeneity because the rims of the platelets would serve as the active sites for exchange. Consequently, the platelet edges would be enriched in Cs, whereas the unreacted cores of the platelets might retain their interlayer Na until the reaction was complete.

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**Dissolution and reprecipitation.** If exchange occurred by complete dissolution of Na-birnessite and re-precipitation of Cs-birnessite, then we might have observed an intermediate...
reaction stage in which diffraction peaks were replaced by a broad amorphous intensity, as was observed by Post et al. (2005) during the dehydration of Mg-, Ba-, and H-birnessite and the subsequent crystallization of hausmannite. However, for all concentrations of the CsCl solutions the diffraction patterns retained their sharpness throughout the exchange processes (Figs. 2 and 9), as was also the case for the K- and Ba-exchanges studied by Lopano et al. (2007). Admittedly, a dissolution stage may have passed too quickly during the Cs exchange to have been captured within our data sampling intervals of two minutes. However, the fact that the Na-birnessite peaks gradually disappeared as the Cs-birnessite peaks emerged argues against total dissolution of Na-birnessite on timescales that were smaller than the temporal resolution of our experiments.

By analogy with the “fine-scale” solution and deposition model of O’Neil (1977), another possibility is that, at any given time-step, only a small amount of end-member Na-birnessite dissolved and re-precipitated as end-member Cs-birnessite. However, in our experiments the lattice parameters of the emergent Cs-rich birnessite-ss phase changed with time, and EDS analysis of batch experiment samples revealed that the transient phases were of intermediate compositions. Therefore, our observations are not consistent with the appearance of a Cs-end-member phase as the Na-end-member dissolved, even on a microscale.

**Delamination and reassembly.** We propose that the exchange of Cs for Na in birnessite occurs through a mechanism that combines aspects of the simple diffusion and the dissolution-reprecipitation models. Specifically, we believe that our results imply a 3-step delamination-reassembly process in which (1) octahedral sheets rift apart from a given interlayer as Cs ions penetrate that interlayer; (2) the Na cations within that interlayer are rapidly replaced wholesale by Cs cations; and (3) the octahedral sheets re-assemble. This mechanism operates on successive layers in a single platelet in a zipper-like sequence, as is shown schematically in Figure 13, although it should be borne in mind that Cs-exchange most likely initiates at multiple interlayers within a single crystal.

This model resolves the contradiction in our XRD and TEM results. As successive interlayers in a single platelet swap out Na for Cs, those individual platelets become biphase mixtures with co-existing Na-rich and Cs-rich layer domains. With increasing time, the Cs-rich layer domains increase in volume at the expense of the initially Na-rich layer domains. X-ray diffraction patterns would reveal separate 00l peaks with increasing exchange, and the peak intensities for the larger d-spacing peak (corresponding to Cs-birnessite) would increase as the smaller d-spacing peaks (Na-birnessite) diminish, just as we observed in our TR-XRD experiments. The change in lattice parameters for the Cs-rich

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**Figure 12.** Comparison of selected-area electron diffraction (SAED) patterns of (a) Na-birnessite, (b) incompletely exchanged Cs,Na-birnessite, and (c) Cs-birnessite grains, taken parallel to the c-axis, in the a-b plane, at a camera distance of 100 cm.

**Figure 13.** Schematic diagram of our proposed delamination-reassembly model for the exchange of Cs for Na in birnessite. When Na-birnessite is immersed in Cs-rich aqueous solutions, the octahedral Mn sheets delaminate successively to allow interlayer Na cations (small circles) to diffuse into the solution as Cs cations (large circles) to migrate into the interlayer region. This model assumes that exchange occurs through wholesale replacements of interlayer regions.
phase may result from the strain created by the sandwiching of Cs interlayers between Na interlayers. As more Cs interlayers are generated, the size of the Cs domains increases and the strain decreases. Over time, the lattice parameters for Cs-birnessite will approach their end-member values.

Our EDS and SAED data are explained by the fact that the crystals we examined occurred as platelets that invariably lay parallel to the supporting foil. Consequently, the electron beam was oriented perpendicular to the octahedral sheets, thereby constraining our sampling direction. With the electron beam parallel to the c-axis, both EDS spectra and SAED patterns included data collected normal to the layers rather than along particular layers. If Cs-substitution occurs via our proposed delamination process, then EDS probes would sample both the Na-rich and the Cs-rich layers, yielding the identical intermediate composition at all points across a given platelet, as was observed. Similarly, electron diffraction patterns of the intermediate compositions would contain information reflective of both Na-rich birnessite (streaking parallel to <110>*) and Cs-rich birnessite (superstructure diffractions at 1/3<310>*), also as was observed.

In essence, our model represents an intermediate pathway between simple diffusion and dissolution-reprecipitation. Unlike a simple diffusion mechanism, our delamination model presumes a step-wise collective substitution of cations, as is consistent with the rapidity of the exchange and the absence of reaction rims. Unlike a dissolution and reprecipitation mechanism, our delamination model maintains the structural integrity of the octahedral scaffolding. We suggest that a delamination-reassembly exchange pathway may be the operative substitution mechanism in other layered structures, perhaps even aluminosilicate clays.

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