

## Ba and Ni speciation in a nodule of binary Mn oxide phase composition from Lake Baikal

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### Abstract

The partitioning and incorporation mechanism of Ni and Ba in a ferromanganese nodule from Lake Baikal were characterized by X-ray microfluorescence, microdiffraction, and absorption spectroscopy. Fe is speciated as goethite, and Mn as romanechite (psilomelane) and 10 Å-vernadite (turbostratic buserite) with minor 7 Å-vernadite (turbostratic birnessite). Barium is associated with romanechite and Ni with vernadite in distinct and irregularly distributed layers, and each type of Mn oxide is separated from the other type by goethite. The binary Mn oxide banding pattern is interpreted by a two-mode accretionary model, in which the variation in Ba flux induced by hydrothermal water pulses determines whether a tectomanganate (romanechite) or phylломanganate (vernadite) is formed during the ferromanganese nodule accretion. Consistent with the dependence of Ni sorption on pH and the circumneutral pH of the lake water, nickel is mainly substituted isomorphically for Mn in the manganese layer, and is not sorbed at vacant Mn layer sites in the interlayer.

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### 1. INTRODUCTION

Lake Baikal in southeastern Siberia is one of the oldest lakes in the world, with an age of around 30 million years. The basin of Lake Baikal belongs to a particular class of rift basins that originate from subsidence along major border faults, resulting in the particular long, deep and narrow shape of the lake trench. It is the world's largest body of fresh surface water, with a maximum depth of 1642 m, and belongs to the category of ultra-fresh calcium bicarbonate water of a pH 7.6, with a dissolved salt content below 100 mg L<sup>-1</sup> (Votintsev, 1961). The concentrations of most elements in the water column are controlled by weathering of surrounding rocks, and the riverine input is consequently

the major source comprising 75% to >99% (97% on average). A major feature that sets Lake Baikal apart from other deep lakes is that its entire water column is constantly enriched with oxygen due to its oligotrophic status (low biological productivity results in a small amount of degradable organic matter), and regular deep-water renewal with oxygen-rich surface waters (Votintsev, 1961; Granina et al., 2000). In the water 2–5 mm above the bottom the oxygen content is never less than 230 μmol L<sup>-1</sup> (Martin et al., 1998). The sediment surface is orange-brown and enriched in Fe and Mn oxyhydroxides, and the oxidized layer is from <1 to 20–30 cm thick, depending on the oxygen penetration depth, and thus on the sedimentation rate (Granina et al., 2000). Another peculiarity of this freshwater system is a sub-aquatic hydrothermal point source (vent system) located in the region of Frolikha Bay (Northern Baikal). The hydrothermal activity is characterized by increased heat flux from the bottom (Crane et al., 1991; Golubev et al., 1993), excess of crustal <sup>4</sup>He in bottom waters (Kipfer et al., 1996),

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peculiarities in chemistry and stable isotope composition in pore waters at the hydrothermal discharge site (Callender and Granina, 1992; Shanks and Callender, 1992; Granina et al., 2002, 2007), and a specific and rather diverse fauna (Gebruk, 1995). Although such hydrothermal manifestations are frequent in extensional tectonic settings, no other vent sites have been found yet in the Baikal rift lake.

Distinct zones of sedimentary Fe and Mn oxyhydroxides are known to occur within the sediments of Lake Baikal (Leybovich, 1983; Granina, 1992a,b; Granina et al., 2000, 2004; Müller et al., 2002). They spread over all three lakes' basins, and also exist on the underwater Academician Ridge that separates the northern and central basins. In many locations extended layers occur in near-surface sediments, with the top part enriched in Mn and the bottom part in Fe. At some locations, solid Fe/Mn crusts up to 1 cm thick are formed at the oxic–anoxic interface. A low sedimentation rate on the order of  $0.01\text{--}1\text{ mm yr}^{-1}$ , and a low content of sedimentary organic carbon of about 2 wt.% on average, is a prerequisite for the post-depositional formation and preservation of such layers and crusts (Granina, 1992b; Granina et al., 2000, 2004). The Mn content in the enriched layers ranges from  $10^{-5}\text{ mol cm}^{-3}$  at sites of higher sedimentation rate near the Selenga Delta, and up to  $10^{-2}\text{ mol cm}^{-3}$  at the Academician Ridge where the sedimentation rate is the lowest and the elements scavenged are of autochthonous origin (hydrogenous accretion mode).

The ferromanganese precipitates have a Fe/Mn ratio of  $\sim 17$  on average in shallow water crusts and concretions, and down to  $\sim 1.4$  in deep-water crusts (Granina, 1992a; Deike et al., 1997). Globe-shaped Mn-rich ferromanganese nodules have been found on the northern slope of Bolshoi Ushkani Island in Northern Baikal during a cruise of submarine "Pisces" in 1990. These nodules had Fe/Mn ratios lower than all other ferromanganese precipitates previously known from the lake, and were enriched in Co, Ni, Cu, and especially Ba (Bukharov et al., 1992; Amirzhanov et al., 1993).

Although ferromanganese precipitates have been studied extensively over the past decades, it is only within the last few years that insights have been gained into the speciation of their trace element inventory (Marcus et al., 2004a, and references therein). In this paper, we examine the speciation of Ni and Ba in one of these Northern Baikal nodules using the combination of synchrotron-based X-ray micro-fluorescence ( $\mu$ -XRF), micro-diffraction ( $\mu$ -XRD), and micro and powder extended X-ray absorption fine structure (EXAFS) spectroscopy (Manceau et al., 2002a). The first technique was used to map Mn, Fe, Ni, and Ba distributions, and to identify elemental associations. Based on these maps, the nature of the minerals hosting Ni and Ba then was deduced from  $\mu$ -XRD and Mn-EXAFS spectroscopy. Finally, with Ni- and Ba-EXAFS spectroscopy, the structural relationship between the trace metals and mineral hosts was examined.

## 2. MATERIALS AND METHODS

### 2.1. Sample

The ferromanganese nodule sample Pa-1 used for this study was described in detail earlier together with another

sample Pa-2 of similar composition (Bukharov et al., 1992; Amirzhanov et al., 1993; Bukharov and Fialkov, 1996). Both samples were collected from the northern slope of the Bolshoi Ushkanyi Island at 510 m depth (Electronic Annex EA-1). The originally subspherical sample Pa-1 of about 2 cm in diameter was encased in epoxy resin and cut in half along its longest axis to obtain cross sections in a plane that included the direction of its highest accretion rate. A thin section of one of the cut surfaces revealed in reflected light a strong texture characterized by zones of botryoidal layering, with individual layers having thicknesses from tens to hundreds of microns. Semi-quantitative electron probe microanalysis (Electronic Annexes EA-2 and EA-3) showed large variations in chemical composition, with about 8–54% MnO, 7–57% Fe<sub>2</sub>O<sub>3</sub>, and up to 6.6% BaO and 1.4% NiO. A self-supporting 30- $\mu\text{m}$  thick micro-polished section was prepared for synchrotron radiation-based X-ray analysis.

### 2.2. X-ray measurements

Micro-XRF,  $\mu$ -XRD, and  $\mu$ -EXAFS measurements were performed on beamline 10.3.2 at the Advanced Light Source (ALS), Berkeley. The beam spot of the microspectrometer can be adjusted from  $16\text{ (H)} \times 7\text{ (V)}$  to  $5 \times 5\text{ }\mu\text{m}^2$  (Manceau et al., 2002a; Marcus et al., 2004b). The distributions of Mn, Fe, Ni and Ba were imaged by scanning the sample under a monochromatic beam and measuring the intensity of the K $\alpha$  (Mn and Ni), K $\beta$  (Fe) and L $\alpha$  (Ba) fluorescence lines with a 7-element Ge solid-state detector and a counting time of 100–150 ms per pixel. The intensities of the K $\alpha$  and K $\beta$  radiation were recorded with an excitation energy of 10 keV, and that of the Ba(L $\alpha$ ) radiation by subtracting the Ti(K $\alpha$ ) intensity measured at 5197 eV (Ba L<sub>3</sub> –50 eV) from the Ba(L $\alpha$ ) + Ti(K $\alpha$ ) intensity at 5297 eV (Ba L<sub>3</sub> +50 eV). The fluorescence yield was normalized against the incident intensity  $I_0$  and the counting time. No attempt was made to calculate elemental concentrations because of the difficulties inherent to this type of quantification, and the focus was on relative element distributions, mineralogy and crystal chemistry of the sample. Elemental distributions were portrayed with a RGB (red, green, blue) color scheme, by assigning two (bicolor map) or three (tricolor map) primary colors to different elements (Manceau et al., 2002a). When fluorescence signals from these elements overlap, the primary colors mix in proportion to the fluorescence intensities and produce a secondary color: green + blue = cyan, green + red = yellow, blue + red = magenta, and red + green + blue = white. The  $\mu$ -XRD patterns were recorded in transmission with a Bruker 6000 CCD binned to  $1024 \times 1024$  pixels at 14 keV incident X-ray energy and  $16 \times 7\text{ }\mu\text{m}$  beam size. The two-dimensional XRD patterns were calibrated with corundum ( $\alpha$ -Al<sub>2</sub>O<sub>3</sub>) and integrated to one-dimensional patterns with the Fit2d code (Hammersley et al., 1996). Mn K-edge, Ni K-edge, and Ba L<sub>3</sub>-edge  $\mu$ -EXAFS spectra were collected at different spots on the sample, representing either Ba- or Ni-rich regions. To avoid possible radiation damage, only one spectrum was collected at each spot, and spectra from nearby ( $\sim 10\text{ }\mu\text{m}$  spacing) spots having similar compositions and

mineralogy, as seen from the XRF maps and  $\mu$ -XRD patterns, were averaged. Several single-scan Ba and Ni spectra were recorded at distant spots throughout the thin section to verify the uniqueness of Ba and Ni speciation. Also, a wide-beam ( $300 \times 150 \mu\text{m}$ ) Ni-EXAFS spectrum was collected on beamline FAME (Proux et al., 2006) at the European synchrotron radiation facility (ESRF, Grenoble, France) on a sample powder. All Ba-EXAFS spectra were statistically indistinguishable, showing that the Ba speciation was unique. The Ni-EXAFS spectra, although distinct from those for Ba, also indicated a single Ni species by their lack of statistically significant variation. EXAFS data were analyzed by standard procedures (Teo, 1986).

### 3. RESULTS

#### 3.1. XRF maps

The alternating Fe-rich (green) and Mn-rich (blue) bands are depicted in the bicolor map of Fig. 1a. When

Ba is added in red, the Mn layers from the lower right region are magenta, some in the middle and upper left region are dark violet, and others from the middle region remain blue (Fig. 1b). In contrast, these blue layers become magenta when Ni is imaged in red (Fig. 1c). Thus, Ba and Ni are associated with Mn, and mostly anti-correlated. A Ba association with Mn rather than with Fe is common in ferromanganese precipitates (Manceau et al., 2007), and was reported in samples from Lake Tomahawk (USA) and Lake Biwa (Japan) (Jones and Bowser, 1978; Sugiyama et al., 1992). The negative relationship between Ba and Ni is imaged in Fig. 1d with Ba in blue and Ni in red. This map contains blue and red layers, but also some magenta and dark violet layers, suggesting an incomplete separation of Ba and Ni. Mineralogical and spectroscopic data presented below show that these two elements are associated with two distinct Mn oxides, which are completely separated in the monochromatic layers and mixed in the bicolored layers from Fig. 1d.

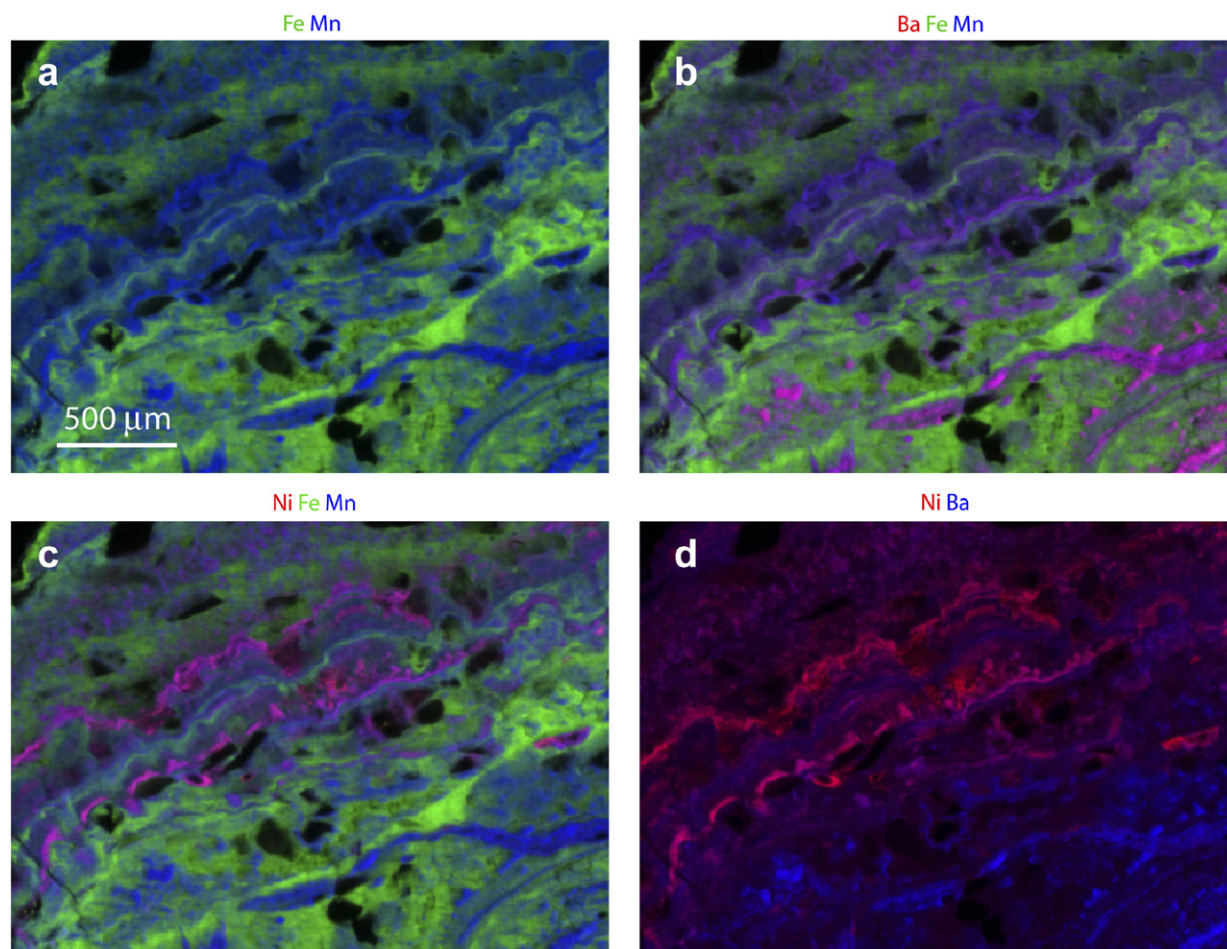


Fig. 1. Micro-XRF fine map of the distribution of Mn, Fe, Ba, and Ni in a ferromanganese nodule from Lake Baikal. The color coding is: (a) green = Fe, blue = Mn; (b) red = Ba, green = Fe, blue = Mn; (c) red = Ni, green = Fe, blue = Mn; (d) red = Ni, blue = Ba. Each pixel is colored in proportion to the normalized fluorescence signals. Size =  $2500 \text{ (H)} \times 1800 \text{ (V)} \mu\text{m}^2$ , pixel resolution and beam size =  $5 \times 5 \mu\text{m}^2$ , counting time = 150 ms/pixel.

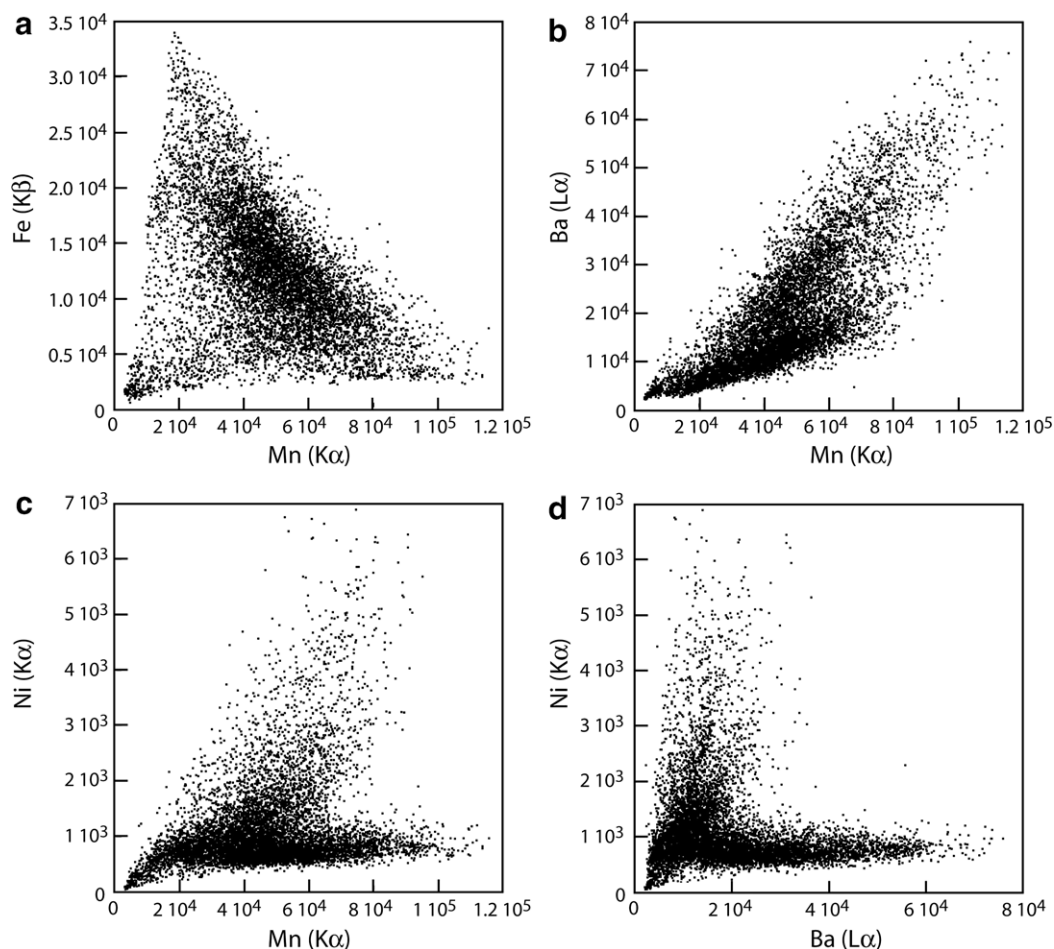
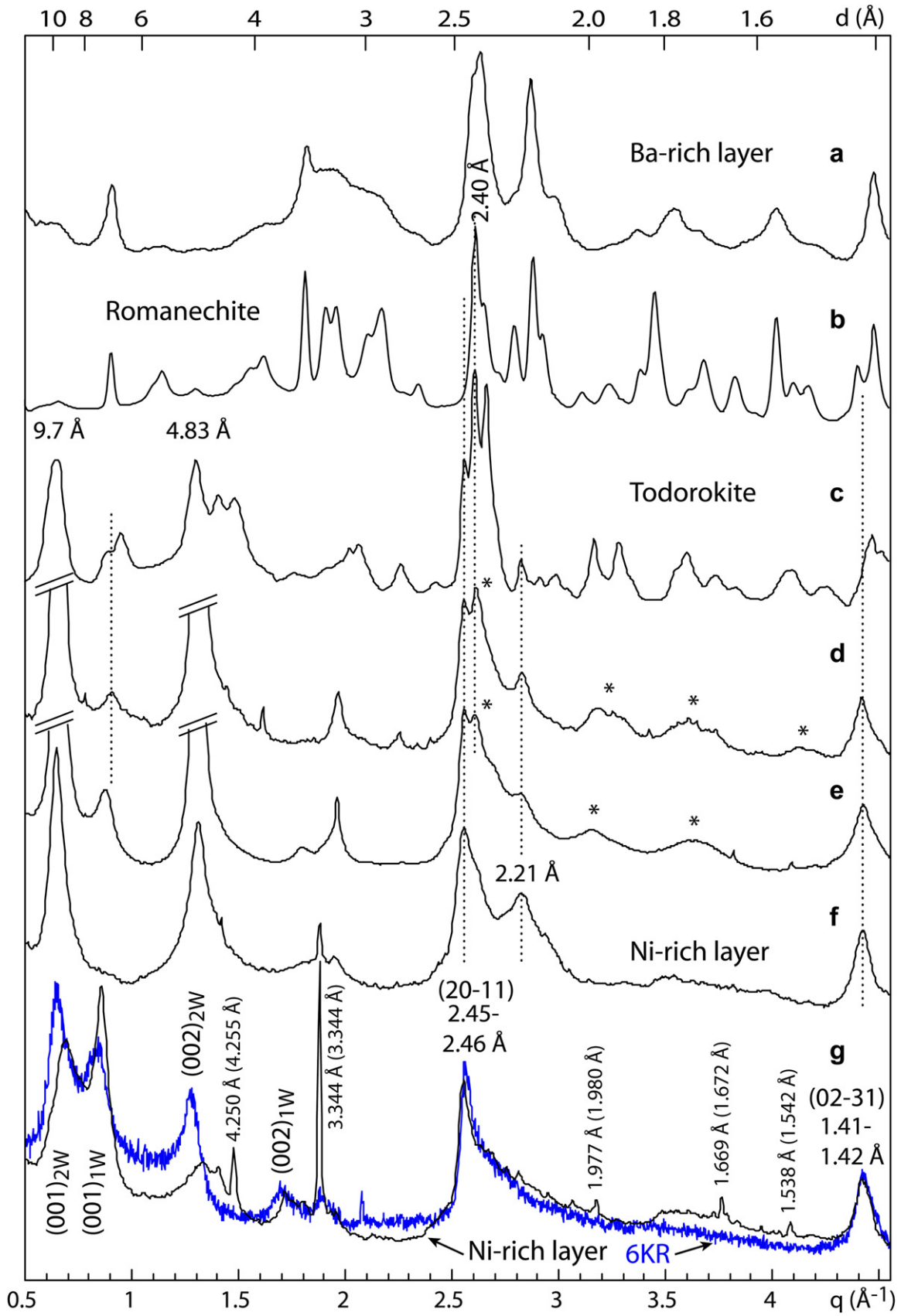


Fig. 2. Scatter plots of the normalized fluorescence counts between Fe and Mn (a), Ba and Mn (b), Ni and Mn (c), and Ni and Ba (d).

Elemental correlations were evaluated in more detail by representing the fluorescence intensities from the previous maps as scatter plots (Fig. 2; Manceau et al., 2002a). The Fe vs. Mn plot reveals that the scanned region contains areas in which the two major elements are well separated, and others which are permeated with both Fe and Mn. The average Fe/Mn ratio is about 1, as estimated from the relative intensities of the Fe(Kβ) and Mn(Kα) signals, correcting the data for air absorption, fluorescence yields, and taking  $I[\text{Fe}(\text{K}\alpha)]/I[\text{Fe}(\text{K}\beta)] = 100/17$ . Chemically, the most differentiated bands contain about 5–10 times more of one major element over the other. The Ba–Mn and Ni–Mn populations are distinguished in the scatter plots of Fig. 2b and c by rays of differing slopes. The Ni–Ba plot

shows mainly two rays along the axes, confirming that where Ba is concentrated, Ni is not, and vice versa. The Ba–Mn scatterplot has a Pearson correlation  $r = 0.95$ , and the Ni–Mn scatterplot  $r = 0.74$ . The variability of the Ba/Mn and Ni/Mn ratios explains the non-uniformity of the magenta hue in Fig. 2b and c. Close examination of the two maps shows that the variations in hue are more pronounced for Ni than for Ba in this region of the sample, in agreement with the lower  $r$  value of the Ni–Mn population. Two other regions of the same ferromanganese nodule were mapped, a large  $4880 \times 3000 \mu\text{m}^2$  area at  $20 \times 20 \mu\text{m}^2$  resolution, and a  $3400 \times 2300 \mu\text{m}^2$  area at  $10 \times 10 \mu\text{m}^2$  resolution. Results are similar to those presented in this section (Electronic Annex EA-4).

Fig. 3. Powder microdiffraction patterns taken at a Ba-rich spot (a) and two Ni-rich spots (f,g), and powder diffraction patterns from references. (b) Romanechite standard from Hufgard quarry, Germany, (c) todorokite standard from Todoroki mine, Japan, (d) hydrothermal ferromanganese oxide from the Costa Rica margin (Bodei et al., 2006), (e) diagenetic marine ferromanganese oxide from Takahashi et al. (2007) (sample coded as 'D465 outer' in this article). 6KR is a natural vernadite (Manceau et al., 2007). Peaks labeled with an asterisk are from todorokite. 1W and 2W denote one- and two-water layer hydrates, respectively. The wavevector  $q$  is defined as  $2\pi/d$ , with  $d$  the  $d$ -spacing. The  $d$ -values of vernadite were calibrated using the X-ray lines of quartz. Theoretical values for quartz are in parenthesis. The accuracy is better than  $0.01 \text{ \AA}$  over the whole  $q$ -range. The two  $hk$  bands at  $2.45\text{--}2.46 \text{ \AA}$  and  $1.41\text{--}1.42 \text{ \AA}$  are indexed by choosing a C-centered layer cell (Villalobos et al., 2006).



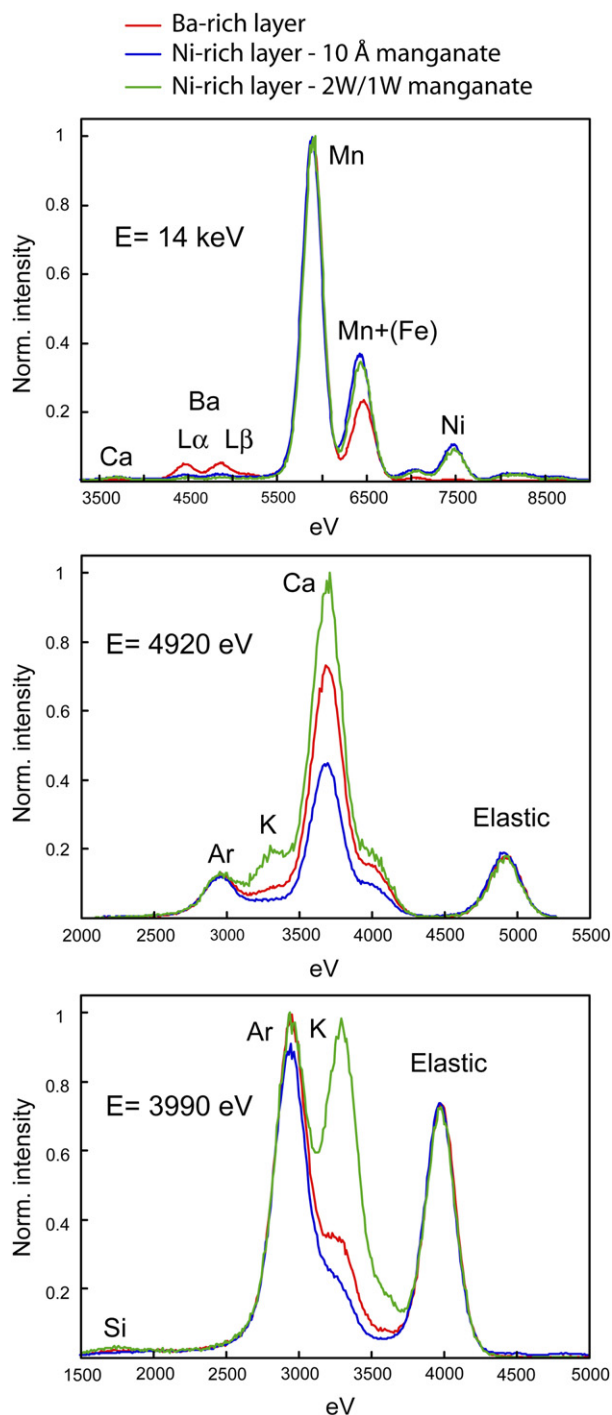


Fig. 4. Characteristic micro-fluorescence spectra of the Ba-rich species and the two Ni-rich species (10 Å and 2W/1W manganates) taken at  $E = 14$  keV (top graph), 4920 eV (i.e., below the Ti K-edge; middle graph), and 3990 eV (i.e., below the Ca K-edge; bottom graph).

### 3.2. X-ray microdiffraction

Two-dimensional  $\mu$ -XRD patterns were collected on 20 spots rich in Fe, Ni, or Ba. All patterns contained continuous rings of constant intensity along the ring perimeters. Since the X-ray beam had a micrometer size, these powder

rings arise from nanometer-sized particles (Manceau et al., 2002a; Electronic Annex EA-5). The diffraction rings from the Fe layers and Fe-rich matrix between the Mn layers matched the  $d$ -spacings for goethite. No other Fe oxide was detected. Some patterns also contained spotty reflections from coarse grains of quartz, feldspar, or mica. The nature of the Mn phases was identified from the radially integrated and background-subtracted one-dimensional patterns (Fig. 3) and the X-ray fluorescence spectra recorded at the same spots as the 2D patterns (Fig. 4).

All patterns from the Ba-rich, Ni-poor layers were similar and, except for the 1.77 Å reflection ( $q = 3.54 \text{ \AA}^{-1}$ ), had rings matching the position of the most prominent powder diffraction lines of romanechite (or psilomelane), a tectomanganate with  $3 \times 2$  tunnel structure (Wadsley, 1953; Turner and Buseck, 1979; Turner and Post, 1988) (Fig. 3a and b). The romanechite specimen from Lake Baikal has smaller crystallites than the reference, as indicated by the broadness of its diffraction peaks. All patterns in Ni-rich, Ba-poor layers look like the one shown in Fig. 3f, with one exception shown in Fig. 3g. The most common patterns have basal diffraction maxima at 9.7 Å (001 reflection) and 4.8 Å (002 reflection), and  $hk$  bands at  $2.455 \pm 0.005 \text{ \AA}$  (20–11 reflection) and  $1.415 \pm 0.005 \text{ \AA}$  (02–31 reflection) that match known diffraction features of 10 Å-vernadite, also named busserite (a discussion on the nomenclature of disordered phyllomanganates can be found in Manceau et al., 2007) (Giovanoli et al., 1970, 1975; Giovanoli and Bürki, 1975; Usui and Mita, 1995). The high intensity of the two basal reflection indicates that vernadite nanoparticles are not intermixed or intergrown with Fe oxide nanoparticles (e.g. ferrihydrite) at the nanoscale, as observed in so-called Fe-vernadite (Burns and Burns, 1979; Manceau and Combes, 1988; Varentsov et al., 1991; Manceau et al., 1992, 2007; Vodyanitskii et al., 2003, 2004). The  $d$ -spacings of the 20–11 and 02–31 reflections are in the ratio of  $\sqrt{3}$ , which indicates that the phyllomanganate layers have hexagonal symmetry with unit cell parameters  $a = b \sim 2.83 \text{ \AA}$  (Drits et al., 1997). The low value of the layer-cell dimension is an indication that the layers do not contain detectable  $\text{Mn}^{3+}$  (Manceau et al., 1997; Gaillot et al., 2007), in agreement with the oxidizing sample environment. As a comparison, synthetic hexagonal birnessite (HBI) equilibrated at pH 4 has 13%  $\text{Mn}^{3+}$  in its layer and  $a$  and  $b$  parameters of 2.848 Å (Lanson et al., 2000). Also, lithiophorite  $(\text{Al}_{0.67}\text{Li}_{0.32})(\text{Mn}_{0.68}^{4+}\text{Mn}_{0.32}^{3+})\text{O}_2(\text{OH})_2$ , which contains 32% of layer  $\text{Mn}^{3+}$  (Yang and Wang, 2003; Manceau et al., 2005), has  $a = b = 2.925 \text{ \AA}$  (Post and Appleman, 1994). The scattering tail of the high- $q$  asymmetrical 20–11 reflection is modulated by a broad reflection at 2.20–2.22 Å. This reflection could not be assigned to any coexisting mineral species and, thus, is part of the phyllomanganate pattern. This reflection has never been observed on synthetic vernadite ( $\delta$ - $\text{MnO}_2$ ), and its structural origin in natural samples has been discussed by Varentsov et al. (1991).

The asymmetrical shape of the 20–11 reflection towards higher  $q$  values is strong evidence that the 10 Å-manganate has a layer structure. Here a parallel can be made with smectites, whose 02–11 band is also asymmetrical on the right side due to the lack of 3D periodicity (Brindley and Brown, 1980; Drits and Tchoubar, 1990). The reflec-

tions near 10 and 2.4–2.3 Å cannot be attributed to todorokite, a tectomanganate with 3 × 3 tunnel structure (Post et al., 2003), because this interpretation is inconsistent with the positions and relative intensities of other reflections. The possibility of a phyllo-manganate–todorokite mixture also can be dismissed from the comparison with marine samples containing the two phases (Figs. 3d and e). Todorokite has a prominent diffraction line at 2.40 Å, seen as a second maximum to the right of the 20–11 reflection for the phyllo-manganate, when the two phases are mixed. Also, the admixture of todorokite modulates the pattern of vernadite between the 20–11 and 02–31 reflections because todorokite has three intense lines in this region, and the phyllo-manganate pattern has none.

The distinct phyllo-manganate pattern recorded at one Ni-rich spot out of ten analyzed by  $\mu$ -XRD has no reflection at 2.20–2.22 Å, and exhibits two sets of 001 and 002 basal reflections, one at 9.1 and 4.7 Å, and another at 7.3 and 3.6 Å, from two-water (2W) layer and one-water (1W) layer diffracting crystallites, respectively. This pattern is extremely similar to that described recently for a vernadite specimen (6KR) formed naturally at the surface of quartz grains from sand filters used in groundwater treatment plants (Manceau et al., 2007). The overlay plot of the two vernadite patterns shows that the two samples do not have the same proportions of 1W and 2W crystallites. Revealing information regarding the average layer composition of each diffracting unit is deduced from the following observations. First, in the new sample the 001 reflection of the 2W component is more asymmetrical and further shifted in the direction of the 1W component than in the quartz coating. Second, the 001 reflection of the 2W component is broader than that of the 1W component. Third, the two basal reflections are irrational for the 2W component (9.1 and 4.7 Å), and rational for the 1W component (7.3 and 3.6 Å). In contrast, the 2W basal reflections have less irrationality (9.7 and 4.9 Å) in the quartz coating.

All these observations, put together, indicate that the 2W crystallites consist of a random interstratification of 2W and 1W layers, while the 1W crystallites contain only, or predominantly, 1W layers (Reynolds, 1989; Manceau et al., 2007). In mixed-layer compounds, the exact position of  $d(001)$  depends on the relative proportions of the two types of layers (Reynolds, 1989). Since the 001 reflection of the 2W component is closer to 9.7 Å than 7.3 Å, the fraction of the 2W layer hydrates is higher than 50%. However, not all 2W diffracting units contain the same proportion of 1W layers, as indicated by the tail of the 001 peak towards smaller  $d$  spacings (i.e., higher  $q$  values). Microfluorescence analysis of 2W–1W vernadite revealed that its interlayer contains K and Ca, whereas only Ca was detected at all other Mn spots

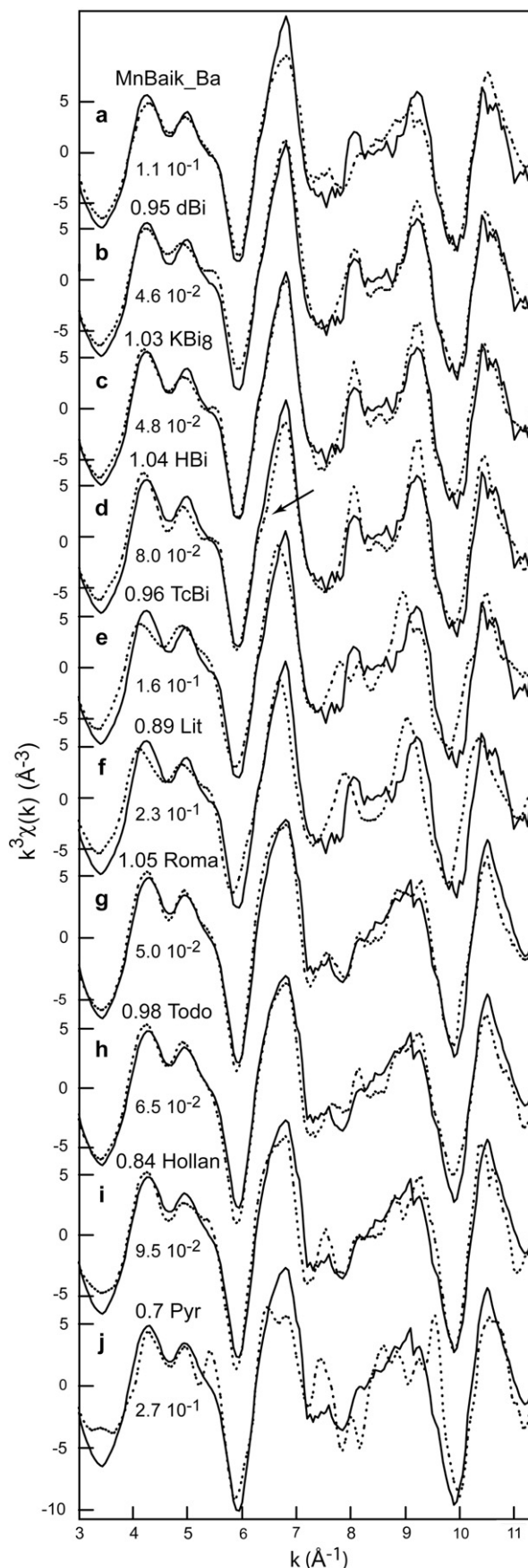


Fig. 5. (a) Micro Mn-EXAFS spectra taken at a Ni-rich spot (MnBaik\_Ni, solid line) and at a Ba-rich spot (MnBaik\_Ba, dotted line). (b–f) Comparison of the MnBaik\_Ni spectrum (solid line) with references (dotted lines). (g–j) Comparison of the MnBaik\_Ba spectrum (solid line) with references (dotted lines). The value attached to each plot is the squares of the residuals, normalized to the sum of the squares of the data values (Norm. Sum-Sq), calculated over the  $[3.0\text{--}11.4 \text{ \AA}^{-1}] k$  interval.

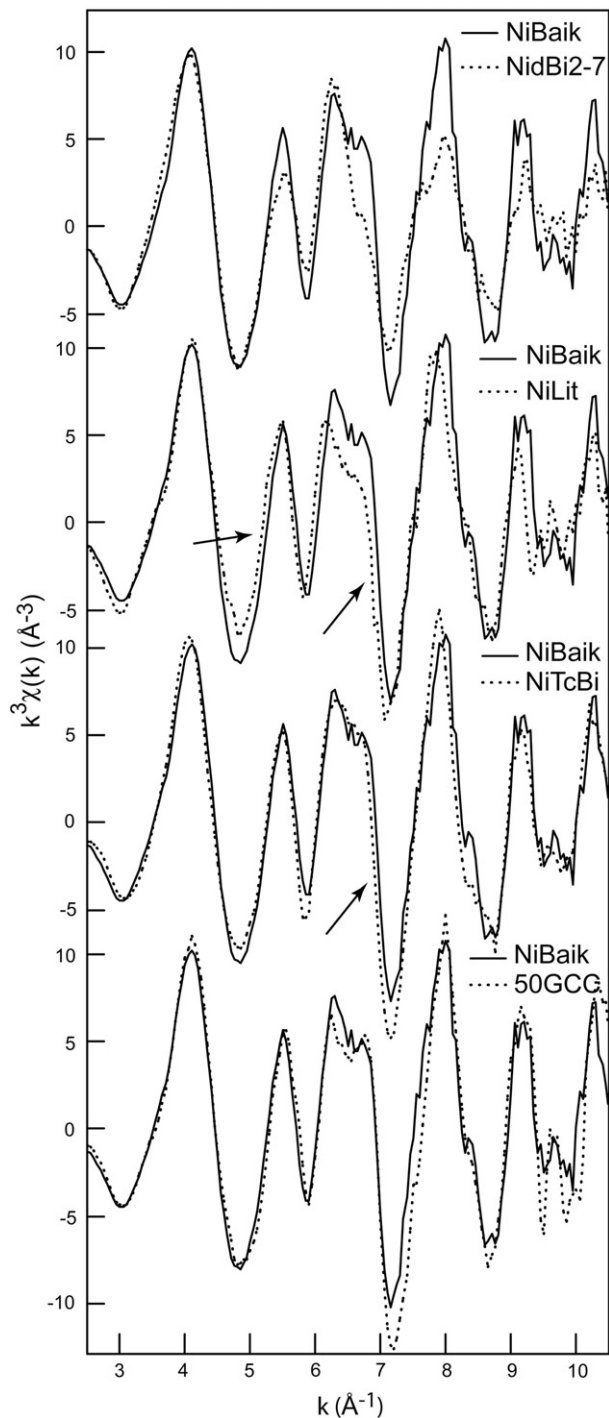


Fig. 6. Ni K-edge EXAFS spectrum of the sample (NiBaik, solid line) and references (dotted lines). Arrows point out the frequency differences which occur when the Mn layer has Mn<sup>3+</sup> (i.e., Ni–Mn<sup>3+</sup> pairs).

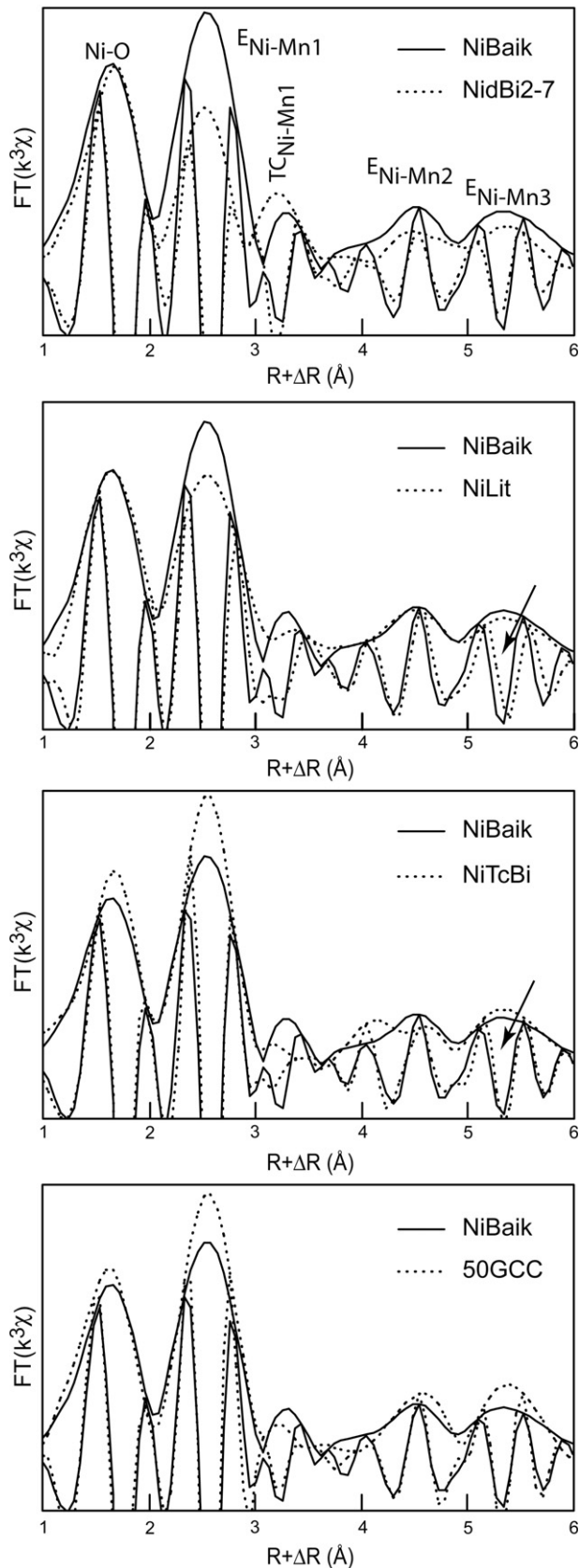


Fig. 7. Fourier transforms (FTs) of the Ni-EXAFS data in Fig. 6. The shift in frequency observed in the [5–7 Å<sup>-1</sup>] *k* interval in Fig. 6 transforms in the real space as a phase difference of the imaginary parts of the Ni–Mn1 and Ni–Mn3 peaks. The phase difference increases with the radial distance.

(Fig. 4). With a cation ionic potential (valency over ionic radius) of only 0.72, compared to 2.0 for Ca, K has a low affinity for water and usually occurs in dehydrated to weakly hydrated interlayers (Ferrage et al., 2005). Consequently, the 1W crystallites in 2W–1W vernadite are probably K-rich and nearly homoionic given the narrowness of their 001 reflection. Also, the proportion of 2W and 1W layers in mixed-layer crystallites should depend on the Ca/K ratio and, thus, no two crystallites are likely to be chemically and structurally identical, in agreement with diffraction data. However, it is not possible to conclude whether the proportion of 2W and 1W layers in a given crystallite varies with the hydration state of the sample because the stability of the 2W layers with relative humidity is unknown. XRD analysis of the bulk sample is inconclusive about this question because the 2W–1W vernadite specimen is minor.

### 3.3. EXAFS spectroscopy

#### 3.3.1. Mn K-edge EXAFS

Mn K-edge EXAFS spectroscopy is sensitive to Mn oxidation state, Mn–O and Mn–Mn interatomic distances, and polyhedral connectivity (Manceau and Combes, 1988). Therefore, it complements the mineralogical and structural information obtained from XRD. Micro-EXAFS spectra were recorded at the same POIs as for  $\mu$ -XRD. The Mn-EXAFS spectra from the Ba-rich and Ni-rich layers are distinct, although each set of spectra from the same chemical type of layer is statistically invariant. The distinction can be seen in the spectral shapes in the  $[7.0\text{--}9.5 \text{ \AA}^{-1}]$  interval and confirms the existence of two separate manganese species (Fig. 5a). There is, however, a similarity in the frequency of all spectra from both species, indicating that the average Mn oxidation state is similar throughout the sample. The  $[7.0\text{--}9.5 \text{ \AA}^{-1}]$  re-

gion, also called the ‘indicator region’ (Marcus et al., 2004a), is sensitive to the amount and ordering of  $\text{Mn}^{4+}$  and  $\text{Mn}^{3+}$  in the manganese layer of phyllosmanganate and the size of the tunnel structure in tectomanganate (Manceau et al., 2002a, 2004, 2005; Gaillot et al., 2003). The form of Mn in the two manganates was determined using spectra of model compounds as structural fingerprints. A selection of candidate species is shown in Fig. 5, and includes a synthetic turbostratic birnessite,  $\text{Na}_{0.24}(\text{H}_2\text{O})_{0.72}(\text{Mn}_{0.94}^{4+}\text{Vac}_{0.06})\text{O}_2$  (dBi, Marcus et al., 2004a; Villalobos et al., 2006); a synthetic hexagonal birnessite,  $\text{K}_{0.231}\text{Mn}_{0.077}^{3+}(\text{Mn}_{0.885}^{4+}\text{Vac}_{0.115})\text{O}_2 \cdot 0.60\text{H}_2\text{O}$ , made from  $\text{KMnO}_4$  (KBi<sub>8</sub>, Gaillot et al., 2003); a synthetic hexagonal birnessite,  $\text{Mn}_{0.05}^{2+}\text{Mn}_{0.12}^{3+}(\text{Mn}_{0.74}^{4+}\text{Mn}_{0.10}^{3+}\text{Vac}_{0.17})\text{O}_{1.7}(\text{OH})_{0.3}$  (HBi), prepared by equilibrating at low pH triclinic Na-birnessite,  $\text{Na}_{0.31}(\text{Mn}_{0.69}^{4+}\text{Mn}_{0.31}^{3+})\text{O}_2 \cdot 0.40\text{H}_2\text{O}$  (TcBi, Silvester et al., 1997); and, a synthetic lithiophorite (Lit, Yang and Wang, 2003). TcBi and Lit have about the same proportion of layer  $\text{Mn}^{3+}$ , but not the same  $\text{Mn}^{4+}$ – $\text{Mn}^{3+}$  cation ordering (Drits et al., 1997; Manceau et al., 2005). The best spectral match to our Mn oxide database of the Ni-containing manganate (MnBaik\_Ni) was obtained with dBi, which has no  $\text{Mn}^{3+}$  and no interlayer Mn (Fig. 5b). In MnBaik\_Ni, vacant octahedral sites (Vac) are capped likely by interlayer Ca since the lake water is rich in calcium bicarbonate, and Ca is difficult to detect as a backscatterer by Mn-EXAFS.

The spectrum of the Mn species from the Ba-rich layers (MnBaik\_Ba) more closely resembles the romanechite spectrum than any other references (Fig. 5g). The MnBaik\_Ba spectrum is clearly different from that of hollandite, a  $2 \times 2$  tunnel structure tectomanganate, and slightly different from that of todorokite. Upon decreasing the size of the tunnel in tectomanganates, the second oscillation broadens and eventually splits into two maxima at 6.5 and 6.8  $\text{\AA}^{-1}$ , as

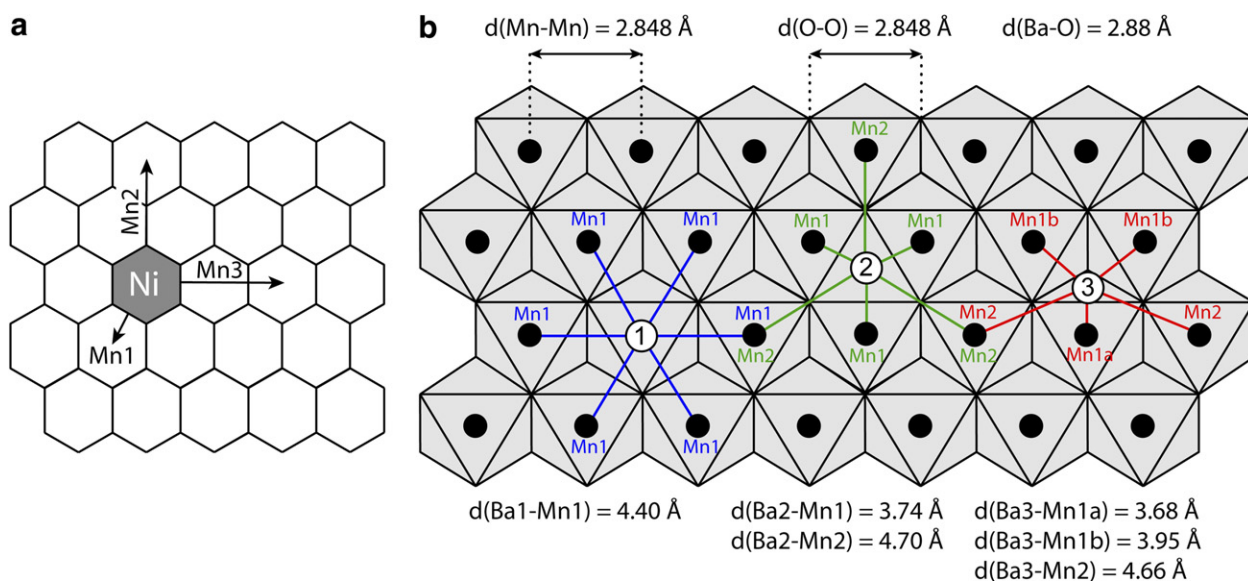


Fig. 8. (a) Representation of the  $^{\text{E}}\text{Ni-Mn1}$ ,  $^{\text{E}}\text{Ni-Mn2}$ , and  $^{\text{E}}\text{Ni-Mn3}$  pairs in a phyllosmanganate layer with  $a = b = 2.848 \text{ \AA}$  (HBi). Theoretical sorption sites of Ba in the interlayer of a phyllosmanganate. At position 1 (Ba1), Ba is above a vacant layer site and is corner-linked to six Mn octahedra from the layer. At position 2 (Ba2), Ba is at the center of a tetrahedral site formed by three edge-sharing Mn octahedra, and is linked to them by corners. At position 3 (Ba3), Ba is shifted from the previous position in direction of one of the three edge-sharing Mn octahedra by  $1/4$  the Ba2–Mn1 distance.

observed in Fig. 5i and j for hollandite and pyrolusite, an  $1 \times 1$  tunnel structure tectomanganate (Manceau and Combes, 1988). Thus, the shape of the second oscillation is sensitive to the proportion of edge- and corner-sharing Mn octahedra in manganates, being symmetrical in the absence of corner-sharing linkage, and otherwise asymmetrical. In phyllosulfates, the width of the second oscillation is indicative of interlayer Mn octahedra linked through corners to layer Mn octahedra (Manceau et al., 1997), as shown in Fig. 5d with HBi which has 17% interlayer Mn. From the comparison of MnBaik<sub>Ni</sub> and KBi<sub>8</sub> (Fig. 5c), the maximum amount of interlayer Mn in the Lake Baikal sample is estimated to be 8%.

### 3.3.2. Ni K-edge EXAFS

Since nickel is always sixfold coordinated in phyllosulfates, the shape and frequency of Ni-EXAFS essentially depend upon the type (e.g., Ni vs. Mn) and position (e.g., layer vs. interlayer) of non-nearest-neighbor atoms, leading to the concept of a ‘structure fingerprint’, as for Mn-EXAFS (Manceau et al., 2007). From the survey of about fifteen reference spectra from layer Ni oxides representing a range of Ni binding environments, four were selected for a demonstration of the fingerprint technique applied to Ni speciation and uniqueness of our structural interpretation (Fig. 6). The Ni<sub>2</sub>Bi<sub>2</sub>-7 reference is a synthetic turbostratic birnessite (dBi) containing a total of 0.002 Ni per Mn atom, of which 45% are substituted for Mn in the layer (<sup>E</sup>Ni complex neighbored by six Mn<sup>4+</sup>), and 55% are in the interlayer at vacant Mn layer site (<sup>TC</sup>Ni complex; Manceau et al., 2007). The NiTcBi reference is a triclinic sodium birnessite (TcBi) doped with 0.0044 Ni per Mn. The NiLit reference is a natural Ni-containing lithiophorite (Manceau et al., 2002b). Finally, the 50GCC reference is a marine Mn<sup>4+</sup> 10Å-vernadite from the Costa Rica margin. This reference has only <sup>E</sup>Ni (Bodei et al., 2006), as do NiTcBi and NiLit.

The Ni-EXAFS spectral shape of the Lake Baikal sample (NiBaik) is similar to those of NiTcBi, NiLit and 50GCC, and distinct from that of Ni<sub>2</sub>Bi<sub>2</sub>-7, suggesting that Ni is predominantly incorporated in the Mn layer of the Baikal sample. The split second oscillation has the same frequency for 50GCC and NiBaik, and is shifted to lower  $k$  values for NiTcBi, and even more so for NiLit. The increase of the EXAFS frequency in the two Mn<sup>3+</sup>-containing references arises because the <sup>E</sup>Ni–Mn<sup>3+</sup> distances are longer than the <sup>E</sup>Ni–Mn<sup>4+</sup> distances. Supporting evidence is seen by comparing the Fourier transforms (FTs), which shows that the imaginary part of the <sup>E</sup>Ni–Mn3 peak, and to a lesser extent of the <sup>E</sup>Ni–Mn1 peak, is shifted to higher distance in NiTcBi and NiLit relative to NiBaik (Fig. 7). The <sup>E</sup>Ni–Mn3 pair is more sensitive to the presence of Mn<sup>3+</sup> around Ni because the distance separation between <sup>E</sup>Ni and Mn3 is twice that of <sup>E</sup>Ni and Mn1 (Fig. 8a). Thus, we conclude that Ni is surrounded only by Mn<sup>4+</sup> atoms in NiBaik, in agreement with the lack of layer Mn<sup>3+</sup> deduced from XRD and Mn-EXAFS data. The FT of NiBaik seems to have a peak at  $R + \Delta R = 3.2\text{--}3.3$  Å, which suggests the occurrence of some <sup>TC</sup>Ni atoms in the interlayer. Data simulations showed that the proportion of <sup>TC</sup>Ni to total Ni is less than 10%, which is consistent with the high amplitude of the <sup>E</sup>Ni–Mn1 peak.

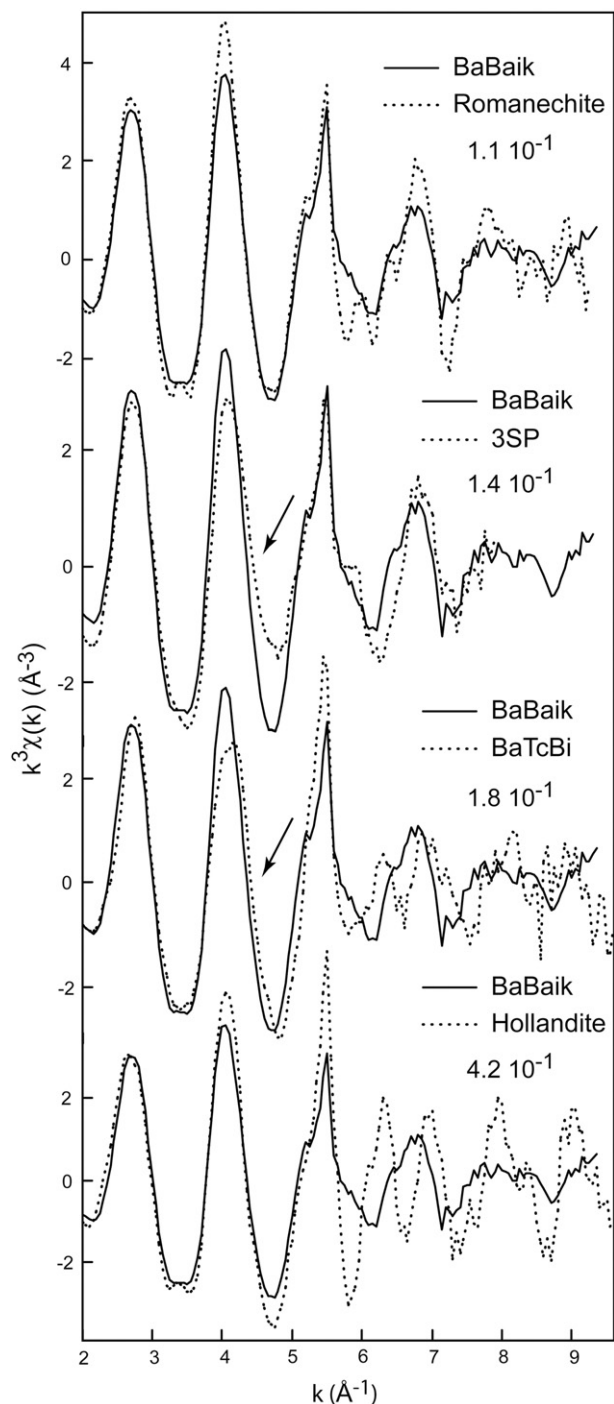


Fig. 9. Ba  $L_3$ -edge EXAFS spectrum of the sample (BaBaik, solid line) and references (dotted lines). The closest match of BaBaik is with romanechite, in agreement with  $\mu$ -XRD data. The sharp peak at  $5.5 \text{ \AA}^{-1}$  is a multi-electronic effect, which does not affect the quantitative analysis and structural interpretation (Manceau et al., 2007). The Norm. Sum-Sq values were calculated over the  $[2.0\text{--}8.0 \text{ \AA}^{-1}] k$  interval.

### 3.3.3. Ba $L_3$ -edge EXAFS

Four references were used to determine Ba speciation: hollandite from Langenberg, Germany (Frondelet et al.,

Table 1  
Results from quantitative analysis of the Ba L<sub>3</sub>-Edge EXAFS data

Sample	Ba–O			Ba–Mn1			Ba–Mn2			$\Delta E$	Res
	<i>R</i> (Å)	CN	$\sigma$ (Å)	<i>R</i> (Å)	CN	$\sigma$ (Å)	<i>R</i> (Å)	CN	$\sigma$ (Å)		
Hollandite <sup>a</sup>	2.90	8.0 <sup>b</sup>	0.13	3.68 4.00	4.0 <sup>b</sup> 8.0 <sup>b</sup>	0.12 0.17	4.66	8.0 <sup>b</sup>	0.13	4.0	9.3
Romanechite	2.89	8.0 <sup>b</sup>	0.13	3.76 3.95	2.0 <sup>b</sup> 4.0 <sup>b</sup>	0.13 0.17	4.70	4.0 <sup>b</sup>	0.13	2.9	13.7
BaTcBi	2.85	6.0 <sup>b</sup>	0.13	3.62 4.08	1.0 <sup>b</sup> 2.0 <sup>b</sup>	0.09 0.16	4.68	2.0 <sup>b</sup>	0.11	1.9	14.1
3SP	2.85	5.3	0.13	3.68 3.92	1.0 <sup>b</sup> 2.0 <sup>b</sup>	0.06 0.10	3.72	2.0 <sup>b</sup>	0.12	2.6	11.2
BaBaik	2.88	8.8	0.16	3.84	1.0	0.13	4.67	1.5	0.13	2.2	14.6

$\Delta E$ , inner potential correction in eV; Res, fit residual defined as  $\sum\{|z_{\text{exp}} - z_{\text{fit}}|\} / \sum\{|z_{\text{exp}}|\}$ ; interval of the inverse Fourier transform: 1.05–3.6 Å;  $sS_0^2 = 1.4$ ; CN and  $\sigma$  values in BaBaik are not individually accurate because these two parameters were highly correlated in the spectral fit.

<sup>a</sup> Crystallographic values are: 8 O at 2.88–2.90 Å, 4 Mn at 3.58–3.65 Å, 8 Mn at 3.99–4.17 Å, and 8 Mn at 4.59–4.64 Å (Miura, 1986).

<sup>b</sup> Fixed value.

1960), romanechite from Hufgard quarry, Germany, Ba-sorbed synthetic TcBi (BaTcBi) and a natural Ba-containing vernadite (3SP). The purity and crystallinity of the three first standards were verified by XRD. The last reference is from quartz coatings, as the 6KR phyllosulfate reference described in the XRD section. In all these references, Ba is located above an empty tetrahedral cavity formed by three edge-sharing MnO<sub>6</sub> octahedra, and off-centered toward one of the three O<sub>layer</sub>–O<sub>layer</sub> edges to an extent that depends on the mineral species (Fig. 8b; Manceau et al., 2007). When the offset in position is important, the first Mn shell is split, which manifests itself in the EXAFS spectrum by a split of the fourth oscillation at 6.0–7.5 Å<sup>-1</sup>, as observed for hollandite and BaTcBi (Fig. 9; Table 1). This oscillation has a single maximum in the two other references and BaBaik, meaning that Ba is closer to the center of the tetrahedral cavity. The exact shape of this oscillation also depends to some extent on the Ba–O distance (or Ba coordination) since the Ba–O wave interferes with the Ba–Mn waves in this spectral region. Thus, care has to be taken when drawing fine structural distinctions based solely on the shape of the fourth oscillation. The distance separation between the two Mn subshells determined by data simulation is 4.08 – 3.62 = 0.46 Å in BaTcBi, 4.00 – 3.68 = 0.32 Å in hollandite, 3.92 – 3.68 = 0.24 Å in 3SP, and 3.95 – 3.76 = 0.19 Å in romanechite (Electronic Annex EA-6). The BaBaik data could be reproduced with an unsplit first Mn shell at 3.84 Å, suggesting that of all manganese species examined here, Ba is the closest to the center of the tetrahedral cavity in BaBaik. The specificity of the Ba environment in BaBaik also is seen clearly on the Fourier transforms (Fig. 10).

The plausibility of our fit models was verified by constructing an idealized MnO<sub>2</sub> octahedral sheet and calculating by geometry alone the Ba–Mn distances for Ba at the center of the tetrahedral cavity (Ba2 position), and shifted in the direction of the nearest Mn shell by 1/4 the Ba2–Mn1 distance (Ba3 position, Fig. 8b). In this model, the unsplit Mn1 shell is at 3.74 Å, and the second-nearest Mn shell (Mn2) at 4.70 Å from Ba2. The split Mn1 shell is at 3.68 and 3.95 Å, and the Mn2 shell at 4.66 Å from Ba3. These

theoretical values are in good agreement with EXAFS distances (Table 1) and support our structural interpretation.

Data simulation also showed that the phase and amplitude of the second EXAFS oscillation are sensitive to Ba coordination, which is right-shifted (i.e., shorter Ba–O distance) and less intense when Ba is sixfold (BaTcBi, 3SP) coordinated than when it is coordinated to eight or more oxygens (hollandite, romanechite; Fig. 9). Using the wave frequency criterion, Ba is at least eight coordinated in the Lake Baikal sample, as confirmed by the similarity of the mean Ba–O distance in BaBaik and romanechite (2.88–2.89 Å; Table 1). The short Ba–O distance (2.85 Å) in phyllosulfate is also an indication that Ba is closer to the Mn layer when the structure is lamellar. Here, the wave amplitude criterion is less conclusive because simulations showed that high structural disorder occurs at the Ba site in BaBaik. In conclusion, Ba is associated with romanechite, but is bonded to a slightly different surface site than in the well-crystallized reference. In addition to the difference in crystallinity, it is possible that the two romanechite specimens do not have the same stoichiometry, with the Lake Baikal species containing less Mn<sup>3+</sup> and more Mn vacancies since the lake sediment–water interface is oxidizing. This hypothesis is supported by the compositional variability of todorokite. Two structural formulae for natural todorokite have been reported in the literature: Mg<sub>0.07</sub><sup>2+</sup>Na<sub>0.06</sub><sup>+</sup>Ca<sub>0.03</sub><sup>2+</sup>K<sub>0.01</sub><sup>+</sup>[Mn<sub>0.80</sub><sup>4+</sup>Mn<sub>0.17</sub><sup>3+</sup>Vac<sub>0.02</sub>Fe<sub>0.01</sub><sup>3+</sup>Al<sub>0.01</sub><sup>3+</sup>]<sub>0.01</sub>O<sub>2</sub>·0.55H<sub>2</sub>O (Gutzmer and Beukes, 2000) and Mg<sub>0.07</sub><sup>2+</sup>Na<sub>0.07</sub><sup>+</sup>Ca<sub>0.02</sub><sup>2+</sup>K<sub>0.02</sub><sup>+</sup>[Mn<sub>0.73</sub><sup>4+</sup>Mn<sub>0.27</sub><sup>3+</sup>]<sub>0.02</sub>O<sub>2</sub>·0.7H<sub>2</sub>O (Post et al., 2003). The specimen analyzed by Gutzmer and Beukes (2000) had a higher Mn<sup>4+</sup>/Mn<sup>3+</sup> ratio and probably contained some vacancies (Vac).

#### 4. DISCUSSION

Dissolved (<0.45 μm) Ni concentrations in waters of Lake Baikal are about 2 nM while riverine inputs are up to 8 nM (Falkner et al., 1997). Nutrient-like vertical profiles and a flow-weighted mean residence time of 110 years, which is one-third of the mean water residence time, indicate an active removal process as a result of biological

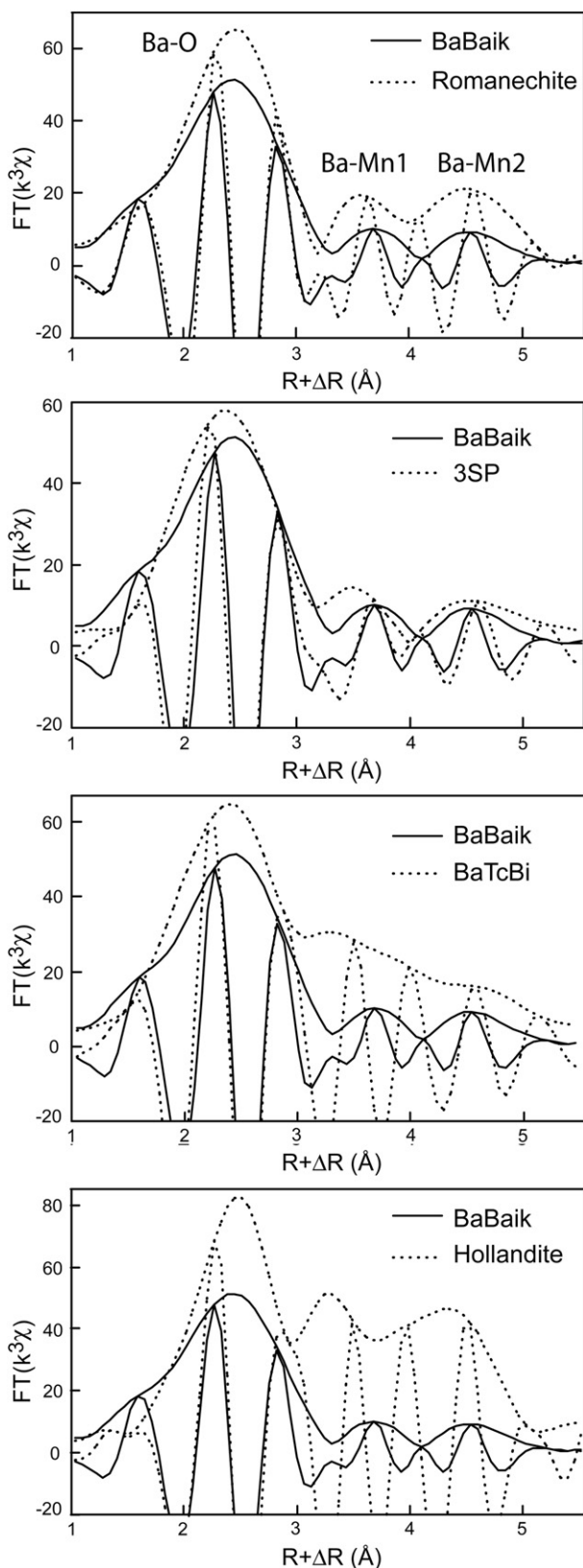


Fig. 10. Fourier transforms of Ba-EXAFS data in Fig. 9. Note the shift to higher  $R + \Delta R$  values of the Ba-Mn1 peak position and phase for BaBaik relative to romanechite.

scavenging. Barium concentrations are about 70 nM (Falkner et al., 1997), comparable to other freshwater lakes with values between 30 and 300 nM (Sugiyama et al., 1992). Since the sulfate concentration is only about 50  $\mu\text{M}$ , the Ba concentration is at a minimum two orders of magnitude undersaturated with respect to the solubility product of  $\text{BaSO}_4$ . Nonetheless, the vertical profile of Ba concentration is slightly depleted (10 nM) near the sediment–water interface, probably as a result of Ba scavenging by Mn nodules (Falkner et al., 1997). The mean flow-weighted residence time of Ba is in the same order as that of water, but a yet unknown amount of Ba may originate also from hydrothermal water discharge, such as that reported at Frolikha Bay, Northern Baikal (Kipfer et al., 1996). Barium concentrations of up to 2  $\mu\text{M}$  controlled by barite solubility were measured in porewaters sampled at Frolikha Bay (Granina et al., 2007).

Although hydrothermal vent activity has not yet been discovered at the B.Ushkanii Island site, some indirect evidence exists. The site is located at crossing faults, and bottom waters from this site have enhanced electrical conductivity (Bukharov and Fialkov, 1996). Moreover, both diverse benthic organisms with new taxa and the hydrothermal precipitate geysierite were observed at this site just as at the Frolikha vent (Takhteev et al., 2001). If present, the hydrothermal activity would be responsible for an enhanced supply of Ba, and its precipitation at some distance from the source as high Ba-containing romanechite. The formation of this mineral is probably driven by the enhanced input of Ba, as this element acts as structure-directing template for the formation of  $2 \times 2$  and  $2 \times 3$  tectomanganates (Giovanoli and Balmer, 1983; Feng et al., 1998). Romanechite forms preferentially to hollandite at ambient condition because the size of the hydrated  $\text{Ba}^{2+}$  ion matches the dimension of the  $2 \times 3$  tunnel of romanechite. In contrast, at higher temperature the  $2 \times 2$  tunnel structure of hollandite is stabilized by anhydrous Ba.

The nickeliferous vernadite is probably not hydrothermal because it would be mixed with romanechite (i.e., their formation would be synchronous). It results more likely from the diagenetic transformation of hydrogenous ferromanganese oxides. A diagenetic rather than hydrogenetic origin is supported by the low Fe/Mn ratio of the Ni-rich layers and the high intensity of the two XRD basal reflections, two characteristics that distinguish chemically and structurally vernadite from Fe-vernadite, the main Mn phase from hydrogenous ferromanganese oxides (Takahashi et al., 2007). According to this interpretation, the romanechite layers would form episodically in response to the activity of hydrothermal plumes superimposed on a steady-state system, in which Ni-rich phyllosilicate layers continuously form by the hydrogenous—early diagenetic process. This scenario is compatible with the existence of Ba-rich, Ni-poor layers because hydrogenetic and diagenetic ferromanganese oxides have a much slower growth rate than hydrothermal oxides (Dymond et al., 1984; Halbach et al., 1988; Takematsu et al., 1989). Therefore, the amount of Ni incorporated from the lake water during the time span of the formation of a Ba-rich hydrothermal layer would be small.

In conclusion, the data support the existence of a sub-aquatic hydrothermal discharge at the nodule sampling spot of B. Ushkanii Island. The intermittent activity of venting episodes on that spot is recorded in the thus analyzed ferromanganese nodule by its particular but not uncommon zoned mineralogy and chemistry. For example, a similar bi-phasic manganate mineral association and Ba enrichment in romanechite has been found recently in shallow submarine sediments vented by diffuse hydrothermal seepage of water at Bahia Concepcion, Baja California Sur, Mexico (Canet et al., 2005). Here, the area of suspicion for venting could be ascertained once other nodules become available. At present, the exploration of the lake-bottom sediment is made difficult by the lack of submarine research logistics at Lake Baikal.

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#### APPENDIX A. SUPPLEMENTARY DATA

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.gca.2007.02.007](https://doi.org/10.1016/j.gca.2007.02.007).

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