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Transformation of Ni-containing birnessite to tectomanganate: Influence and fate of weakly bound Ni(II) species

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Abstract

The geochemical behavior of nickel, an essential trace metal element, strongly depends on its interactions with Mn oxides. Interactions between the phyllomanganate birnessite and sorbed or structurally incorporated Ni have been extensively documented together with the fate of Ni along the transformation of these layered species to tunnel Mn oxides (tectomanganates). By contrast, interactions of phyllomanganates with weakly bound Ni species [hydrated Ni, Ni (hydr)oxides], that possibly prevail in natural Ni-rich (>10% NiO) manganates, have received little attention and the influence of these Ni species on the phyllomanganate-to-tectomanganate transformation remains essentially unknown. A set of phyllomanganate precursors with contrasting contents of Ni was thus prepared and subjected to a reflux treatment mimicking the natural phyllomanganate-totectomanganate conversion. Layered precursors and reflux products were characterized with a combination of diffractometric, spectroscopic, thermal, and chemical methods. Ni is essentially present as hydrated Ni(II) and Ni(II) (hydr)oxides in Ni-rich layered precursors whereas Ni(II) sorbed at particle edges prevail at low Ni content. No Ni sorbed at layer vacancy sites or structurally incorporated was detected in the initial vacancy-free layered precursors. Consistent with the high content ($\approx 1/3$) of Jahn-Teller distorted Mn(III) octahedra in layered precursors, which is favorable to their conversion to tectomanganates, Ni-free samples fully convert to an *a*-disordered todorokite, a common tectomanganate with a 3×3 tunnel structure. Contrastingly and despite similar high Mn(III) contents in Ni-rich precursors, hydrolysis of interlayer Ni²⁺ and polymerization of Ni(OH)₂ in phyllomanganate interlayers is kinetically favored during reflux process. Asbolane, a phyllomanganate with an incomplete - island-like - octahedral layer of metal (hydr)oxides, is thus formed rather than todorokite. A nitric acid treatment, aiming at the dissolution of the island-like interlayer Ni(OH)2 layer, allows an easy and unambiguous differentiation between asbolane and todorokite, the latter being unaffected by the treatment. Both compounds exhibit indeed similar interplanar periodicities and can be confused when using X-ray diffraction, despite contrasting intensity ratios. Migration rate of Mn(III) out of the MnO₂ layer relative to the metal hydrolysis and polymerization rate determines the formation of todorokite or asbolane. Here, Ni (OH)₂ polymerization hampers the formation of tectomanganates and likely contributes to the prevalence of phyllomanganates over tectomanganates in natural Ni-rich environments. Most Ni is retained during the reflux process, part of Ni ($\approx 20\%$) being likely structurally incorporated in the reaction products, thus enhancing the sequestration of Ni in Mn oxides. © 2019 Elsevier Ltd. All rights reserved.

Keywords: Nickel; Buserite; Asbolane; Todorokite; Mn oxides

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1. INTRODUCTION

Phyllomanganates and tectomanganates are the two mineral families accounting for most Mn oxides found in terrestrial and aquatic environments (Post, 1999; Peacock and Sherman, 2007a). In the former, edge-sharing MnO_6 octahedra form layers whose symmetry depends on the abundance and distribution of Mn(III) octahedra, whereas in the latter, chains of edge-sharing MnO₆ octahedra form the walls or floor/ceiling of tunnels. Birnessite is, together with its disordered structural analogue vernadite, the most common phyllomanganate and consists of an octahedral layer, ideally MnO₂, whose layer charge deficit originates from the presence of heterovalent layer Mn(II,III) cations and/or of layer vacancy sites. This deficit is compensated for by the presence of cations sorbed at layer vacancy sites or present as interlayer hydrated species (7.0-7.2 Å layerto-layer distance). Heterovalent Mn cations and/or foreign transition metal cations can be hosted in both families, either structurally substituted for Mn(IV) or as charge compensating cations located in phyllomanganate interlayers or in tectomanganate tunnels (Post, 1999). The occurrence of natural phyllomanganates as micro- to nano-sized cryptocrystalline minerals usually enhances their surface reactivity that originates from their high negative layer charge and from the coexistence of heterovalent Mn cations in these structures. As a result, natural phyllomanganates are highly reactive with respect to both organic (Remucal and Ginder-Vogel, 2014) and inorganic compounds. Specifically, natural phyllomanganates are readily associated with transition metals such as Co (Chukhrov, 1982; Manceau et al., 1986, 1987, 1992a, b; Kay et al., 2001; Takahashi et al., 2007; Zawadzki et al., 2018), Zn (Marcus et al., 2004; Manceau et al., 2007b; Lanson et al., 2008; Spinks et al., 2017), or Ni (Chukhrov, 1982; Manceau et al., 1987, 1992a, b, 2002b, 2003, 2007a, b, 2014; Marcus et al., 2004; Bodeï et al., 2007; Peacock and Sherman, 2007a; Roque-Rosell et al., 2010; Dublet et al., 2012; Hein and Koschinsky, 2014; Spinks et al., 2017; Ploquin et al., 2018; Zawadzki et al., 2018), rare earth (Kasten et al., 1998; Kuhn et al., 1998; Ohta and Kawabe, 2001; Takahashi et al., 2007; Hein and Koschinsky, 2014; Ohnuki et al., 2015; Zawadzki et al., 2018), and radioactive elements (Duff et al., 1999; Post, 1999).

In an effort to disentangle the mobility, (bio)availability, and fate of these elements in different geological settings. structural interactions of phyllomanganates with such foreign elements, especially with transition metals, have thus attracted the attention of the scientific community over the last few decades. In most studies, foreign elements are structurally substituted for Mn(IV) in the Mn octahedral layer (e.g. Co - Manceau et al., 1997; Yin et al., 2011a, 2011b, 2015; Simanova and Peña, 2015) or sorbed as inner-sphere complexes either at vacant layer sites and/or at particle edges (e.g. Zn - Lanson et al., 2002b; Manceau et al., 2002a, 2007b; Marcus et al., 2004; Toner et al., 2006; Kwon et al., 2009; Grangeon et al., 2012; Yu et al., 2013; Simanova and Peña, 2015; Qin et al., 2018). Similar to other transition metals, presence of Ni(II) forming inner-sphere complexes species at vacancy or edge sites

and/or structurally incorporated in phyllomanganate layers has been extensively documented (Manceau et al., 2007a, b; Peacock and Sherman, 2007a, b; Grangeon et al., 2008; Peña et al., 2010; Yin et al., 2012, 2014; Kwon et al., 2013; Simanova et al., 2015). Under low pH conditions, Ni(II) occurs mainly as triple- or double-corner sharing (TCS and DCS, respectively) inner-sphere complex above/below vacancy or at edge sites of the phyllomanganate layers (Peacock and Sherman, 2007b; Grangeon et al., 2008; Peacock, 2009; Peña et al., 2010; Simanova et al., 2015). Migration from TCS/DCS sites to octahedral layer sites is favored by increasing contact time and increasing pH (Peacock, 2009), this structural incorporation in octahedral MnO₂ layers being considered as reversible (Peacock, 2009; Peña et al., 2010) and essentially restricted to low Ni contents (Bodeï et al., 2007; Peacock and Sherman, 2007a, b; Kwon et al., 2013; Simanova et al., 2015; Atkins et al., 2016), consistent with natural Ni-poor occurrences (NiO $\approx 0.1-2$ wt.%, - Manceau et al., 2002b, 2007a, b; Bodeï et al., 2007; Peacock and Sherman, 2007a).

At higher Ni contents, hydrated Ni(II) (Turner and Buseck, 1981; Peacock and Sherman, 2007a) and Ni(II) (hydr)oxides (Chukhrov et al., 1987; Manceau et al., 1987, 1992a, b) may also be associated with Mn oxides in a weaker way compared to inner sphere complexation and structural incorporation. The presence of hydrated Ni (II) species in phyllomanganate interlayers leads to the presence of two planes of interlayer water molecules (buserite), as reported by Manceau et al. (2007b). At high Ni contents, Ni(II) (hydr)oxides may also polymerize as complete or incomplete brucite-like octahedral sheets in phyllomanganate interlayers, leading to lithiophorite- or asbolanelike species, respectively. In all cases, the layer-to-layer distance increases from ≈ 7.2 Å (birnessite or vernadite) to \approx 9.4–10.2 Å (buserite, asbolane, and lithiophorite – Burns et al., 1983; Chukhrov et al., 1987; Golden et al., 1987; Manceau et al., 1987, 1992a, b, 1997, 2014; Feng et al., 2001; Bodeï et al., 2007; Wegorzewski et al., 2015). Despite their widespread occurrences in Ni-rich natural environments (NiO \approx 10–20 wt.% – Llorca, 1988; Roque-Rosell et al., 2010; Ploquin et al., 2018), these weaker associations have received little attention compared to structurally incorporated Ni or to Ni inner-sphere complexes. In particular, the stability of these weakly associated Ni species and their possible mobilization upon structural transformation of their host Mn oxides remain poorly documented. On the other hand, the presence of hydrated Ni(II) in phyllomanganate interlayers has been described as favorable to the transformation of phyllomanganates to tectomanganates upon hydrothermal treatment (Ching et al., 1999; Luo et al., 1999; Onda et al., 2007), similar to Mg(II) (Golden et al., 1986, 1987). In this case, todorokite, a common tectomanganate with a uniform 3×3 tunnel structure and intense diffraction lines at 9.6 Å and 4.8 Å (ICDD #38-0475 - Post and Bish, 1988), may be obtained.

The present article thus focuses on the association of weakly bound Ni(II) species with phyllomanganates. Synthetic samples with contrasting Ni contents were thus prepared and characterized by X-ray diffraction, thermogravimetric analysis, and extended X-ray absorption fine structure spectroscopy (Mn and Ni K-edges). In addition, phyllomanganate samples were hydrothermally treated and reaction products characterized to investigate the effect of Ni on the phyllomanganate-to-tectomanganate transformation (Golden et al., 1986; Shen et al., 1993; Feng et al., 2004, 2010; Atkins et al., 2014; Grangeon et al., 2014, 2015; Zhao et al., 2015) and the fate of weakly associated Ni(II) species during this transformation. The negative impact of Ni, when present at high content, on this transformation is compared to that of Co (Wu et al., 2019). As described above, buserite-, lithiophorite-, and asbolanelike phyllomanganates all have interplanar periodicities similar to those of todorokite, however, leading to their frequent confusion in identification (Burns et al., 1983). Special care was thus paid to differentiate these species using nitric acid treatment (Wu et al., 2019) and anomalous Xray diffraction. An optimum procedure allowing this differentiation is proposed, and the present study provides new insights into the interactions between Ni species and Mn oxides in natural Ni-rich environments, thus shedding light on the influence of phyllo- and tectomanganates (asbolane and todorokite, respectively) on Ni-cycling in these contexts. More generally, factors responsible for the transformation of birnessite/buserite to todorokite or asbolane are inferred.

2. MATERIALS AND METHODS

2.1. Synthesis of layered precursors and reflux products

Both initial layered precursors and reflux products described in the present study were prepared as in the previous work on their Co analogues (Wu et al., 2019). Briefly, layered precursors were prepared after Feng et al. (2004) and Song et al. (2010) by adding a concentrated NaOH solution into a Ni(II) + Mn(II) solution prepared from their chloride salts. The resulting solution was then mixed and bubbled with oxygen gas for 5 h. Resulting black precipitates were washed and one half was dried (60 °C, 3 days) for subsequent analyses. Initial Ni / (Ni + Mn)ratios were 0.00, 0.01, 0.05, 0.10, 0.15, and 0.20. Accordingly, initial layered samples were named Bir, NiB1, NiB5, NiB10, NiB15 and NiB20, and these Ni-containing samples are collectively referred to as NiB. The other half of the precipitates was used to tentatively prepare todorokite through a reflux process following the protocol of Feng et al. (2004). Wet precipitates were thus exchanged with Mg²⁺, rinsed, re-suspended in deionized water, and finally refluxed at 100 °C for 24 h. Reflux products were filtered, washed, and dried (60 °C, 3 days). Transformation products of Bir, NiB1, NiB5, NiB10, NiB15 and NiB20 are named Tod, NiT1, NiT5, NiT10, NiT15 and NiT20, respectively; they are collectively referred to as NiT except for Tod.

2.2. Characterization of layered precursors and reflux products

The elemental composition of all samples was determined in triplicate using atomic absorption spectrometry (Varian AAS 240FS) from 0.1000 g of sample dissolved in 25 mL solution of 0.25 mol L^{-1} NH₂OH HCl and 5 mL of 1 mol L^{-1} H₂SO₄. In addition, the specific surface area of the reflux products was determined with the Brunauer-Emmett-Teller (BET) method and a standard adsorption analyzer (Quantachrome Autosorb-1, JEDL-6390/LV). TGA was performed on a NETZSCH TG 209 instrument with a heating rate of $10 \,^{\circ}\mathrm{C}\,\mathrm{min}^{-1}$ and a N₂ flow of 20 mL min⁻¹. Finally, a non-reducing nitric acid treatment (Wu et al., 2019) was performed to differentiate tectomanganates (todorokite) from phyllomanganates (buserite-, lithiophorite, or asbolane-like phases – Wu et al., 2019). Briefly, 0.1 g sample was put in 250 mL of a 1 M HNO₃ solution with soft stirring at ambient temperature (20 °C). At different time intervals over a 1 week period, 5 mL of the suspension was collected and readily filtered through a 0.22 µm membrane to determine Mn, Ni, and Mg release. After 1 week of this HNO3 treatment, residual samples were dried and analyzed with X-ray diffraction (XRD). Comparison of XRD patterns before and after this nitric acid treatment allows differentiating asbolane and todorokite (Wu et al., 2019). In the case of asbolane, dissolution of the interlayer brucite-like layer is induced by the treatment and leads to a collapse of the layer-to-layer distance to the ≈ 7.2 Å periodicity, whereas periodicities are maintained for todorokite, the tectomanganate framework being unaltered, and positions of XRD maxima are thus unchanged.

Mineralogy of all samples was determined from powder XRD analysis using a Bruker D8 Advance diffractometer equipped with Cu K α source ($\lambda = 0.15418$ nm) and operated at 40 kV and 40 mA. XRD patterns were collected at $0.02^{\circ} 2\theta$ intervals using a continuous scan mode. XRD data was collected on fresh and acid-treated samples (all dried at 60 °C for three days) at a scan speed of 10° 2θ min⁻¹. To assess the stability of their hydration, XRD data was collected also on fresh samples after heating to 140 °C. Finally, XRD patterns were collected at 1° $2\theta \min^{-1}$ for aged samples that were kept dry in the dark for three months. Synchrotron XRD patterns were also collected on selected reflux products on beamline BL14B1 at the Shanghai Synchrotron Radiation Facility (SSRF) and on beamline 4B9A at the Beijing Synchrotron Radiation Facility (BSRF). At BSRF, XRD data was collected at the Mn K-edge (6.5 keV), Ni K-edge (8.3 keV), and 12 keV with a typical beam size of $2 \times 1 \text{ mm}^2$, whereas the 18 keV data was collected at SSRF with a typical beam size of $\approx 0.2 \times 0.3 \text{ mm}^2$. In both cases, X-rays were monochromatized with a double-crystal Si (111) monochromator. Finally, high-energy X-ray scattering data was collected at beamline 11-ID-B at the Advanced Photon Source (Argonne National Laboratory, Argonne, IL), using a \approx 58.66 keV X-ray energy ($\lambda = 0.2114$ Å). Pair distribution functions (PDF) and differential PDFs were calculated as described elsewhere (Qiu et al., 2004; Li et al., 2011).

X-ray absorption spectra were collected at the Mn and Ni K-edges on the 1W1B beamline of BSRF to determine the relative proportions of Mn(II), Mn(III), and Mn(IV) species and the local environments of Mn and Ni in both layered precursors and reflux products. Data was recorded at room temperature in transmission (Mn) and fluorescence (Ni) modes, and the energy was calibrated using metallic Mn/Ni foils as references. Extended X-ray absorption fine structure (EXAFS) spectra were processed using Athena (Ravel and Newville, 2005). X-ray absorption near-edge structure (XANES) spectra of both layered precursors and reflux products were used to determine Mn average oxidation state (AOS) with a specific linear combination fitting method (the Combo method). Mn(II), Mn(III), and Mn(IV) reference compounds were those of the original study (Table 1 in Manceau et al., 2012). Linear combination fitting of Ni K-edge EXAFS data was performed with Athena over the 3–10.5 Å⁻¹ k-range whereas the path simulation was performed using Artemis from 1.12-3.4 Å in r-space and over 2.8–11.6 Å⁻¹ in k-space.

3. RESULTS

3.1. Elemental composition

With increasing Ni content, the Mn content decreases from \approx 55.4 \pm 1.0 to \approx 44.8 \pm 1.8 wt.% for layered precursors and from \approx 54.4 \pm 2.1 to \approx 49.2 \pm 0.9 wt.% for reflux products (Table 1). The Ni/(Mn + Ni) molar ratios are essentially similar in layered precursors and their respective reflux products, with only a very limited fraction of Ni being lost either during the Mg-for-(Na, Ni) exchange or during the subsequent transformation process. In addition, the Mg wt.% in the reflux products significantly decreases from $\approx 3.95 \pm 0.08$ wt.% in the Ni-free Tod sample to \approx 2.11 wt.% in NiT20, thus suggesting a competition between Mg and Ni for interlayer and/or tunnel sites of layered precursors and/or of reflux products. Specific surface area (SSA) of NiT samples, as measured with the BET method, is systematically higher than that of Ni-free Tod, but no systematic evolution of SSA is observed with the increase of Ni content.

3.2. Powder X-ray diffraction

XRD patterns collected on fresh Bir (not shown) are essentially similar to those recorded after a few months of aging and display reflections at 7.2, 3.6, \approx 2.45, and \approx 1.42 Å, typical for birnessite (Drits et al., 1997; Lanson

et al., 2002a). Similar reflections are observed for fresh NiB5 and NiB10 samples (Fig. S1), in addition to a broad maximum peaking at ≈ 4.6 Å. The intensity of this maximum increases with increasing Ni concentration in the initial solution and is thus likely related to the presence of Ni (hydr)oxide (ICDD #13-0229), whose formation was reported under similar experimental conditions (oxidation of a Ni-rich alkaline solution - Feitknecht et al., 1956). Consistently, the intensity of this maximum is increased further for NiB15 and NiB20. These two samples exhibit also reflections at ≈ 2.45 , and ≈ 1.42 Å, typical for layered manganates, the basal reflections of which indicate a 9.9–10.0 Å layer-to-layer distance (Fig. S1). Upon moderate heating (140 °C), these reflections shift to indicate a 7.2 Å layerto-layer distance typical for birnessite, as the result of interlayer cation partial dehydration (Golden et al., 1986; Manceau et al., 1987; Drits et al., 1997). The initial presence of two planes of interlayer H₂O molecules in NiB15 and NiB20 (buserite) is most likely indicative of the presence of interlayer hydrated Ni(II) cations, whose hydration is reduced upon heating. Sample heating also results in a decrease of the intensity of the \approx 4.6 Å maximum.

After aging for about three months, all layered precursors but NiB20 display over the low-angle region $(5-30^{\circ} 2\theta)$ strong basal reflections at \approx 7.2 and \approx 3.6 Å typical for birnessite and indicative of the spontaneous dehydration of NiB15 interlayers (Fig. 1). Two planes of H₂O molecules are still present in most NiB20 interlayers (buserite), however. Splitting of reflections at \approx 2.45 and \approx 1.42 Å, and more generally positions of the reflections over the high-angle region (30–85° 2 θ – Fig. 1), unambiguously indicate that initial Bir is a triclinic birnessite (ICDD #43-1456), consistent with the synthesis protocol (Lanson et al., 2002a). With increasing Ni content, high-angle reflections broaden significantly thus indicating a decrease in the coherent scattering domain size. The broad maximum at \approx 4.6 Å is present also in aged samples.

Reflux products systematically show reflections at ≈ 9.6 Å, ≈ 4.8 Å, ≈ 3.2 Å, ≈ 2.46 Å, ≈ 2.39 Å, ≈ 2.22 Å, ≈ 1.95 Å, ≈ 1.73 Å, ≈ 1.52 Å, and $\approx 1.42-1.41$ Å, that are typical of todorokite (ICDD #38–0475 – Atkins et al., 2014, 2016). For the Ni-free Tod, relative intensity of the three low-angle reflections (at ≈ 9.6 , ≈ 4.8 , and ≈ 3.2 Å)

Table 1

Ch	emi	cal	composition	and	specifi	c surfa	ice area	(SSA	1)	of	layered	l precursors	and	refl	ux	produ	icts.
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Samples	Mn	Ni	Ni/(Mn + Ni)	Mg	(Mg + Ni)/Mn	SSA (m^2/g)
I	wt. %	wt. %	mol%	wt. %	mol%	(, e)
Bir	55.4(1.0)	_	_	_	_	_
NiB1	54.1(1.1)	0.39(1)	0.67	_	_	_
NiB5	53.7(1.0)	1.86(4)	3.18	_	_	_
NiB10	51.2(1.0)	3.89(7)	5.98	_	_	_
NiB15	47.3(1.7)	5.32(3)	8.71	_	_	_
NiB20	44.8(1.8)	7.14(7)	11.50	_	_	_
Tod	54.4(2.1)	-	_	3.95(8)	16.4	90.7
NiT1	54.3(8)	0.33(1)	0.57	3.51(12)	16.2	_
NiT5	54.2(6)	1.57(2)	2.57	3.12(2)	15.6	93.7
NiT10	52.7(6)	3.79(1)	5.94	2.61(2)	17.5	119.7
NiT15	52.4(1.2)	5.66(9)	8.41	2.59(6)	20.4	109.0
NiT20	49.2(9)	7.74(2)	11.38	2.11(2)	22.5	106.8



Fig. 1. XRD patterns of layered precursors (top) and reflux products (bottom). Colors correspond to the Ni/(Ni + Mn) ratio of the samples. Green ticks at the bottom of upper and lower figures indicate the reflections of triclinic birnessite (ICDD #43-1456) and todorokite (ICDD #38-0475), respectively.

decreases in the order $I_{9.6} > I_{4.8} \gg I_{3.2}$, consistent with relative intensities of todorokite 001/100, 002/200, and 003/300 reflections (Post et al., 2003), respectively. With increasing Ni content, the positions of both \approx 9.6 and \approx 4.8 Å reflections shift towards higher angles for NiT15 and NiT20, compared to Ni-free Tod (Fig. S2) and the \approx 4.8 Å reflection becomes predominant (Fig. 1), consistent with previous reports of todorokite prepared in the presence of various divalent metals (Shen et al., 1993, 1994; Ching et al., 1999; Nicolas-Tolentino et al., 1999; Fuertes et al., 2012). Although not described as such in these references, this intensity ratio modification is consistent with the presence of asbolane, a phyllomanganate with an incomplete – island-like – octahedral layer of metal (hydr)oxides in the interlayer (Chukhrov et al., 1987; Golden et al., 1987; Fan and Gerson, 2015). The presence of these incomplete brucite-like octahedral layers between MnO_2 layers precludes the collapse of the layer-to-layer distance to ≈ 7.2 Å upon heating (Fig. S3). Note however that positions of the diffraction peaks from tunnel structures such as todorokite are not affected by such a thermal treatment.

3.3. Anomalous XRD patterns

To investigate further the possible origin of the observed contrasting $I_{4.8}/I_{9.6}$ intensity ratios, XRD patterns of Tod, NiT15, and NiT20 were recorded at different energies, including Mn and Ni K-edges to modify the contribution of these elements to the structure factor (F_{hkl}). These XRD patterns logically exhibit reflections at similar

d-values (Fig. 2a-d). Except for the patterns recorded at the Mn K-edge all Tod patterns display similar low $I_{4.8}/I_{9.6}$ intensity ratios (Fig. S4), consistent with previous reports (Shen et al., 2011). Contrastingly, the $I_{4.8}/I_{9.6}$ intensity ratio is significantly higher for NiT15 and NiT20 compared to Tod, and varies significantly for these two samples as a function of the energy used for data collection. Specifically, NiT15 and NiT20 display highest $I_{4.8}/I_{9.6}$ intensity ratios at the Mn K-edge and lowest ratios at the Ni K-edge, whereas similar intermediate ratios are obtained for XRD patterns recorded at energies different from the two absorption edges (Fig. 2a-d; S4). Both features indicate contrasting structures of Tod and Ni-rich NiT products.

As reported in the early descriptions of asbolane, this compound exhibits high values of the $I_{4.8}/I_{9.6}$ intensity ratio (Chukhrov, 1982; Llorca, 1988; Fan and Gerson, 2015),

consistent with XRD patterns calculated for such a structure hosting an incomplete Ni(OH)2 layer sandwiched in between two MnO₂ layers (Fig. 2e). In addition, for a given composition, this ratio increases when the contribution of Mn atoms to the structure factor is reduced (XRD data recorded at the Mn K-edge - Fig. 2a) and decreases when the contribution of Ni atoms to the structure factor is reduced (XRD data recorded at the Ni K-edge - Fig. 2b), consistent with the data. Consistently, this ratio increases also with the completeness of interlayer plane (Fig. 2e). Together with the collapse of laver-to-laver distance upon acid-dissolution of the incomplete Ni(OH)2 layer (see below), these variations of the $I_{4.8}/I_{9.6}$ intensity ratio can be ascribed to the presence of asbolane, rather than todorokite, in the reflux products of Ni-rich precursors (NiT15 and NiT20).



Fig. 2. XRD patterns collected for NiT20, NiT15, and Tod at (a) Mn K-edge (6.5 keV), (b) Ni K-edge (8.3 eV), (c) 12.0 keV, and (d) 18.0 keV. Intensities were systematically normalized to the most intense reflection. Colors as in Fig. 1. (e) XRD patterns calculated for an asbolane containing 0.525 Ni(OH)₂ in MnO₂ interlayers. Contents of Ni and Mn atoms are reduced by 20% at their respective absorption edges to simulate the anomalous effect. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

3.4. Nitric acid treatment

Upon equilibration in nitric acid, Mn release from layered precursors reaches equilibrium within 12 hrs. The proportion of Mn released from NiB is higher than that released from Bir (Fig. 3a), possibly as an effect of reduced crystal size. For reflux products, the release of Mn is slower, reaching equilibrium after \approx 48 hrs, and is essentially the same for all NiT samples (Fig. 3b), except for NiT5. In all cases, Mn release remains limited (0.10–0.18 and 0.13– 0.15 for NiB and NiT samples, respectively), thus indicating a marginal dissolution of the mineral Mn framework. Ni release is both slower and much larger than that of Mn, reaching a plateau (\approx 1.0 and \approx 0.8 for NiB and NiT samples, respectively) after \approx 168 hrs in both layered precursors (Fig. 3c) and reflux products (Fig. 3d). The proportion of Ni released differs however in the two series of samples as essentially all Ni is released from layered precursors whereas $\approx 20\%$ of Ni remains in the reflux products, possibly indicative of Ni structural incorporation. Finally, Mg is systematically totally released from reflux products, possibly at a slower rate with increasing Ni content (Fig. 3f).

After acid treatment, the XRD pattern of Tod is essentially alike that of the initial reflux product of Bir and indicates the sole presence of todorokite in the reflux product, as expected for the hydrothermal transformation of Mn^{3+} -rich phyllomanganate precursors (Fig. 3e). On the other hand, XRD patterns recorded on NiT15 and NiT20 after acid treatment exhibit, over the low-angle region, peaks at \approx 7.2 and \approx 3.6 Å, typical for the phyllomanganate birnessite, rather than peaks at \approx 9.6, \approx 4.8, and \approx 3.2 Å, as expected for todorokite (Fig. 3e). These two XRD patterns



Fig. 3. Proportion of metal cations released during the nitric acid treatment: (a-b): Mn, (c-d): Ni, and (f): Mg. The amount of metal released is normalized to the sample overall metal content (see text for details). (e): XRD patterns of samples treated with nitric acid. Colors as in Fig. 1 except for Bir in (e). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

are actually similar to the one recorded on Bir after acid treatment and match XRD pattern of hexagonal birnessite (ICDD #23-1239), which was obtained from the low-pH equilibration of an initial triclinic birnessite (Lanson et al., 2000). The evolution of XRD patterns recorded from reflux products upon acid treatment is consistent with that observed for an asbolane-like phyllomanganate (Wu et al., 2019) thus indicating the sole presence of this Mn oxide in NiT15 and NiT20. The structure of this ≈ 9.6 Å phyllomanganate consists of an octahedral MnO₂ layer similar to that of birnessite, with an incomplete octahedral Me(OH)₂, MeOOH, or MeO₂ sheet, possibly incommensurate with the MnO₂ layer, in the interlayer (Chukhrov, 1982; Manceau et al., 1987; Fan and Gerson, 2015). This incomplete interlayer octahedral sheet is solubilized by the acid treatment leading to the observed 7.2 Å layer-to-layer distance (Wu et al., 2019). Such a peak at ≈ 7.2 Å is visible in the XRD pattern of acid-treated NiT5 and NiT10 together with a peak at ≈ 9.6 Å, and its intensity increases with increasing Ni content. This dual feature indicates the coexistence of both todorokite-like and asbolane-like reaction products in NiT5 and NiT10. With increasing Ni content, the peak at ≈ 7.2 Å becomes the sole visible diffraction signature in acid-treated NiT15 and NiT20 that contain only asbolane-like phyllomanganate (Fig. 3e).

3.5. Thermogravimetric analysis

Thermogravimetric analyses of layered precursors systematically show a significant weight loss at 140-150 °C corresponding to the dehydration of interlayer Na⁺ and Ni²⁺ cations, this weight loss decreasing with increasing Ni content (Fig. 4a, c; Table 2). A minor weight loss (2.4-2.9% in Bir, NiB5, NiB10, and NiB15) is also observed for these samples below 100 °C, which is most likely related to the departure of adsorbed water (Gaillot et al., 2003). This low-temperature weight loss is significantly increased for NiB20 (9.25%), however, most likely as the result of a low-temperature buserite-to-birnessite transition associated with the loss of one plane of interlayer H₂O molecules as observed during moderate heating of fresh NiB samples (Fig. S1). A significant weight loss (4.10 and 6.72% for NiB5 and NiB20, respectively) is observed also for NiB samples over the 170-400 °C range. This weight loss is essentially absent in the TGA curve of Bir (1.78%) and increases with Ni content. It is likely related to the presence of the Ni (hydr)oxide responsible for the broad XRD maximum peaking at ≈ 4.6 Å (Fig. 1). Finally, a weight loss is observed at \approx 550 °C for all layered precursors, with no significant influence of the Ni content, probably indicating the reduction of MnO₂ to Mn₂O₃ (Bish and Post, 1989).



Fig. 4. Thermogravimetric analysis traces obtained for (a) layered precursors and (b) reflux products, and their 1^{st} derivatives (c and d, respectively). 1^{st} derivative curves were smoothed using a Savitzky-Golay filter and a 15 points window. Colors as in Fig. 1. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Table 2 Weight loss of layered precursors and reflux products over specific temperature intervals.

Temperature (°C)	Weight loss/%							
	Bir	NiB5	NiB10	NiB15	NiB20			
RT-100	2.45	2.87	2.64	2.84	9.25			
100-170	8.43	7.14	6.54	5.73	4.87			
170-400	1.78	4.10	4.74	5.78	6.72			
Total	17.38	19.20	18.85	19.15	25.31			
	Tod	NiT5	NiT10	NiT15	NiT20			
RT-110	2.75	2.93	3.39	3.90	4.12			
110-240	8.05	7.68	7.43	6.57	6.12			
240-380	5.47	5.32	5.04	5.34	5.97			
380-500	2.69	2.33	2.08	1.93	1.79			
500-650	4.50	4.35	4.06	3.75	3.61			
Total	23.47	22.61	22.01	21.48	21.60			

TGA curves recorded for reflux products significantly differ from those of their layered precursors, with weight losses at ≈ 150 , ≈ 285 , ≈ 440 , and $\approx 545 \,^{\circ}\text{C}$ for Tod (Fig. 4b, d). The weight loss observed from ≈ 110 to 240 °C is most likely related to the loss of the H₂O molecules hydrating cations present in todorokite tunnels (Bish and Post, 1989). With increasing Ni content, this weight loss decreases (from 8.05 to 6.12% in Tod and NiT20, respectively) and shifts towards lower temperatures, most likely indicative of the mineralogical evolution evidenced by XRD data recorded on acid-treated reflux products (Fig. 3e). The low-temperature weight loss in NiT15 and NiT20 likely corresponds indeed to the departure of H₂O molecules coexisting with island-like fragments of interlayer octahedral sheets, as reported in hydroxy-interlayer phyllosilicates (Lanson et al., 2015). For NiT10, this lowtemperature weight loss splits in a doublet owing to the coexistence of todorokite and asbolane (Fig. 3e). The weight loss occurring at \approx 285 °C is also ascribed to the departure of tunnel water (Bish and Post, 1989). TGA curves obtained for Tod and NiT20 also differ significantly over this range, with a shift of the weight loss towards higher temperatures in NiT20 compared to Tod. Evolution with increasing Ni content, is not as systematic as that observed for the lowtemperature weight loss, however (Fig. 4d). The intensity of a third weight loss occurring at \approx 440 °C decreases with increasing Ni content to vanish for NiT15 and NiT20 which essentially contain asbolane (Fig. 3e). A last weight loss occurs at 520-550 °C in all reflux products, a temperature similar to that reported for layered precursors and attributed to the reduction of Mn(IV) (Fig. 4b, d).

3.6. X-ray absorption spectroscopy

3.6.1. Mn K-edge XANES

Relative proportions of Mn(II), Mn(III), and Mn(IV) species and Mn average oxidation state (AOS) have been determined using the Combo method (Table 3 – Manceau et al., 2012). Results consistently indicate high ($\approx 1/3$) relative proportions of Mn(III) in layered precursors, which is considered a key condition for the phyllomanganate-

Table 3

Fractional and average valence states of Mn obtained for layered precursors and reflux products from the Combo fit of XANES spectra and 1st derivatives. The estimated error for Combo method is $\pm 4\%$ (Manceau et al., 2012; Yin et al., 2015).

Sample	Mn^{2+} at. %	Mn^{3+} at. %	Mn^{4+} at. %	Mn-AOS
Bir	3	26	72	3.69
NiB1	2	25	74	3.71
NiB5	2	30	67	3.64
NiB10	2	31	68	3.66
NiB15	0	31	69	3.69
NiB20	0	35	65	3.65
Tod	2	25	74	3.72
NiT1	1	35	65	3.64
NiT5	3	32	66	3.63
NiT10	2	35	64	3.62
NiT15	1	34	65	3.64
NiT20	0	33	67	3.67

to-tectomanganate conversion (Cui et al., 2008; Atkins et al., 2014, 2016;Grangeon et al., 2014, 2015; Zhao et al., 2015; Li et al., 2016). The measured proportion of Mn (III) is slightly lower in Ni-free samples compared to the Ni-bearing ones (Table 3) and no significant difference is observed between layered precursors and their reflux products. In all cases, the Mn(II) content is null within the experimental error.

3.6.2. Mn K-edge EXAFS

EXAFS spectra of both layered precursors and reflux products show significant evolutions with increasing Ni content, thus indicating that the presence of Ni affects the local structure of manganates in the two series of samples (Fig. 5). The EXAFS spectrum collected for Ni-free layered precursor (Bir) is typical for triclinic birnessite over both the staircase and the indicator regions $(4-6 \text{ \AA}^{-1} \text{ and}$ \sim 8–9.4 Å⁻¹, respectively – Marcus et al., 2004). In particular, the splitting of the features at ≈ 7.8 Å⁻¹ and ≈ 9.0 Å⁻¹ (arrows in Fig. 5a) is indicative of a high Mn(III) content, of the systematic elongation of Mn(III) octahedra along the *a* axis, and of their ordered distribution in rows parallel to the b axis (Marcus et al., 2004; Gaillot et al., 2007), typical features for triclinic birnessite (Marcus et al., 2004; Webb et al., 2005; Gaillot et al., 2007; Yin et al., 2014). With increasing Ni content, the $\approx 7.8 \text{ Å}^{-1}$ feature becomes less pronounced even though its position remains unchanged, consistent with the similar Mn(III) contents of all layered precursors (Table 3) and with the similar overall frequency of the EXAFS spectra of these samples (Gaillot et al., 2007). The steady lowering of the \approx 7.8 Å⁻¹ feature resolution is thus most likely related either to disorder in the distribution of elongated Mn(III) octahedral or to some randomness in the direction of the elongation. The split $\approx 9.0 \text{ Å}^{-1}$ feature observed for Bir and NiB1 also turns into a broad and more symmetrical feature with increasing Ni content and the induced vanishing of the sharp high-frequency contribution.

In k-space Mn K-edge EXAFS spectra of Tod and NiT1 exhibit sharp features at ≈ 8.0 and ≈ 9.2 Å⁻¹ that are typical for todorokite (Webb et al., 2005; Bodeï et al., 2007; Feng



Fig. 5. Mn K-edge EXAFS spectra obtained for (a) layered precursors and (b) reflux products. Colors as in Fig. 1. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

et al., 2010; Atkins et al., 2014). These two features essentially decrease in intensity at higher Ni content. In addition, the $\approx 9.2 \text{ Å}^{-1}$ feature shifts to lower wave numbers in NiT samples compared to Tod and NiT1 (Fig. 5b). EXAFS spectra of Ni-rich NiT samples and their Fourier transforms (FTs) are actually alike those of their corresponding layered precursors (Fig. 6a and b), consistent with the similar structure of the MnO₂ layer in both birnessite and asbolane. On the contrary, spectra obtained for Ni-free Bir and Tod differ significantly, as expected for layered and tunnel manganese oxides (Fig. 6c). In particular, the FTs of these spectra show a clear shortening of the average Mn-Me_E distance in Tod compared to Bir, as shown by the shift of the imaginary part of the FT (Fig. 6d, arrow), while Mn-Me_E distance remains stable in Ni-rich NiT samples (Fig. 6b). In addition, Tod FT shows a clear maximum at R $+ \Delta R \approx 3.0$ Å, typical of Mn-Me_C pairs, which is not observed for its Bir precursor. For both Ni-free and Nirich samples, Mn-O distances are similar in both layered precursors and corresponding reflux products, consistent with their similar Mn AOS.

Relative proportions of layered and tunnel Mn oxides in reflux products were assessed from the linear combination fitting (LCF) of their Mn K-edge spectra using the corresponding NiB and Tod spectra as standard references for layered and tunnel structures, respectively. Results from the LCF results (Fig. S5; Table 4) confirm the decreasing proportion of tunnel structures in reflux products with increasing Ni content, consistent with XRD data recorded after nitric acid treatment.

3.6.3. Ni K-edge EXAFS

EXAFS spectra collected at the Ni K-edge are essentially similar for both NiB samples, on the one hand, and NiT, on the other hand (Fig. 7), thus implying similar Ni speciation in each of the two sample groups. The feature from 7 to 9 Å^{-1} in the spectra of NiB samples changes however with increasing Ni content. This feature appears as a broad flat maximum in NiB5; with increasing Ni content its intensity decreases towards increasing k values and a minor sharp maximum appears on its high-k end (black arrows in Fig. 7a). Spectra from the two



Fig. 6. Comparison of Mn K-edge EXAFS (a, c) and of the corresponding FTs (showing both modulus and imaginary parts – b, d) obtained for Ni-rich (Top: NiB15 and NiT15) and Ni-free (bottom: Bir and Tod) layered precursors and corresponding reflux products.

Table 4	
Proportions of phyllomanganate and tectomanganate	in reflux
products derived from linear combination fitting of Mn	K-edge χ
(k).	

Sample	Phyllo. (NiBn)	Tecto. (Tod)	Error
NiT5	0.62	0.38	0.06
NiT10	0.83	0.17	0.05
NiT15	0.88	0.13	0.01
NiT20	0.89	0.11	0.02

groups of samples significantly differ from those of Bir- $Ni_{V/S}$ in both k- and r-spaces (Fig. 7), thus implying minimal contribution of Ni associated to hexagonal birnessite either incorporated in the Mn octahedral layer (Bir-Nis) and/or sorbed at layer vacancy sites (Bir-Niv - Atkins et al., 2016). These spectra rather exhibit similarities with those of Ni associated with birnessite/δ-MnO2 varieties containing significant amounts of Mn(III) within the layers (Fig. 7a - Peacock and Sherman, 2007b; Simanova et al., 2015). In both species (Mn^{III}-δ-MnO₂ and TcBi), Ni is essentially sorbed at particles edges, most likely as the result of the limited content of vacant layer sites, sharing either edges or corners with Mn octahedra (Simanova et al., 2015). Consistently, a satisfactory fit to the spectra of NiB5 was obtained as a LCF of Ni associated to such Mn(III)-rich phyllomanganates ($\approx 90\%$), of aqueous Ni, and of Ni in Ni oxide (Fig. 7b; Table 5). Modifications affecting spectra at the Ni K-edge when increasing Ni content could not be satisfactorily reproduced with the same LCF approach, most likely because no reference of the Ni (hydr)oxide identified with XRD (ICDD #13–0229) was available. Consistently, shell-by-shell fitting of the EXAFS data indicates the prevailing contributions of Ni-Ni pairs from NiO and Ni(OH)₂ in Ni-rich layered precursors (Fig. S6, Table S1). In addition to the precipitation of this Ni (hydr)oxide, increasing Ni contents lead to the decrease of crystal size deduced from XRD peak broadening, and to the increased contribution of aqueous Ni, as shown by the higher hydration state of NiB15 and NiB20 compared to NiB5 and NiB10 (Figs. 1, S1).

The systematic comparison of NiT and NiB EXAFS spectra in k-space reveals similar features between the two groups of samples, indicative of a minor evolution of Ni speciation during reflux. NiT spectra exhibit however a minor decrease in amplitude at ≈ 3.8 , ≈ 5.2 , ≈ 6.3 and ${\approx}7.6~{\rm \AA}^{-1}$ and a subtle shift to higher frequencies compared to their NiB counterparts (red arrows in Fig. 7a). Although it was not possible to obtain a decent fit to this data using LCF of available Ni reference spectra, the evolution of Ni speciation during the reflux process was assessed by fitting EXAFS spectra of reflux products as a linear combination of their respective NiB precursors and of Ni reference spectra (Fig. 7b; Table 5). For all NiT samples, a good fit was obtained with $\approx 15\%$ of a Ni(OH)₂ contribution in addition to the layered precursor (Table 5), consistent with the polymerization of a partial Ni(OH)2 layer in birnessite interlayers to form an asbolane-like structure.



Fig. 7. Comparison of Ni K-edge EXAFS spectra obtained for layered precursors and reflux products (black and red solid lines, respectively) with reference spectra (a) and linear combination fitting of Ni K-edge $\chi(k)$ for NiB5 and NiT (b – see text and Table 5 for details on individual contributions to these fits). Bir-Ni_v and Bir-Ni_s represent Ni sorbed above/below vacancy sites and structurally incorporated into MnO₂ layers, repectively (Atkins et al., 2016); Ni(II)–H₂O represents aqueous Ni(II); Ni- δ MnO2 and Ni-TcBir represent Ni sorbed to Mn(III)-rich vernadite and birnessite, respectively (Simanova et al., 2015). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Table 5
Relative proportions of the different contributions to Ni K-edge spectra of NiB5 and of reflux products determined from linear combination
fitting (Fig. 7b).

Samples	Ni-aq	Ni-δMnO ₂	Ni-TcBir	NiO	NiBn	Ni(OH) ₂
NiB5	7(5)	65(6)	23(8)	5(1)	_	_
NiT5	_	_	_	-	81(2)	19(2)
NiT10	_	_	_	_	86(1)	14(1)
NiT15	_	_	_	_	87(1)	13(1)
NiT20	_	_	_	_	88 (1)	12(1)

"-" indicates that the reference was not used during the linear combination fitting.

Ni-TcBir and Ni- δ MnO₂ references correspond to TcBi pH8 (Ni 0.02) and to MnIII_ δ MnO₂ (Ni 0.01) from Simanova et al. (2015), respectively.



Fig. 8. Pair distribution functions obtained for Ni-rich (NiT10 and NiT20) and Ni-free (Tod) reflux products (top). Middle: d-PDFs relative to Tod PDF data; bottom: PDFs calculated for Ni (hydr)oxides (ICSD #76670 and #28101 for NiO and Ni(OH)₂, respectively). Intensities of experimental PDFs and of difference PDFs are normalized to the 2.9–3.1 Å peak.

3.7. Pair distribution functions (G(r))

Atomic PDFs computed from Tod, NiT10, and NiT20 high-energy X-ray scattering patterns show similar peak positions (Fig. 8), indicative of similar local structures (Zhu et al., 2012) as expected from the major contribution of layers/ribbons of edge-sharing MnO₆ octahedra in both layered precursors and reflux products. Consistently, the maxima at ≈ 1.90 Å, ≈ 2.90 Å, and ≈ 3.50 Å essentially correspond to the first neighbor Mn-O, to edge-sharing Mn-Me, and to a combination of corner-sharing Mn-Me and of second neighbor Mn-O atomic pairs, respectively (Zhu et al., 2012). The increased intensity of the ≈ 3.50 Å maximum and the decreased intensity of the one at ≈ 5.00 Å in the PDF of Tod compared to NiT10 and NiT20 are typical of the prevalence of tunnel structures in the former sample and of layered structures in the latter (Grangeon et al., 2015). The \approx 5.00 Å maximum is mainly related indeed to second neighbor Mn-Mn pairs (Figs. 8, S7 - Zhu et al., 2012), that are favored by the lateral extension of MnO_2 layers. Finally, the peak at ≈ 4.50 Å is most likely related to Mn2-Mn2 and Mn4-Mn4 pairs across "holes" in tunnel walls, floors, and ceilings (Fig. S7), consistent with the increased intensity of this ≈ 4.50 Å peak with decreasing tunnel size (Zhu et al., 2012).

Differential PDF (d-PDF) were calculated to assess the specific modifications affecting the reflux products, and more especially the broadening of the \approx 1.90 and 2.90 Å peaks towards high d-values with increasing Ni content (Fig. 8). The d-PDF curves obtained for NiT10 and NiT20, both show a first peak at \approx 2.05 Å typical for Ni-O pairs (Manceau et al., 2007a, b; Peacock and Sherman, 2007a, b; Yin et al., 2012; Kwon et al., 2013; Simanova et al., 2015; Atkins et al., 2016). A second peak, similar in both d-PDF curves, is visible at \approx 3.05 Å. This second peak is intermediate between those calculated Ni-Ni pairs in NiO and Ni(OH)₂, consistent with the presence of both compounds identified in the reflux products with XRD (Figs. 1 and 2).

4. DISCUSSION

4.1. Structural transformation of layered precursors

Addition of Ni during the synthesis of layered precursors does not impact significantly their crystal structure. In particular, the Mn(III) content of layered precursors remains about constant at $\approx 1/3$ (Table 3) whatever the initial Ni content. Consistently, XRD results (Fig. 1) show that layer symmetry remains orthogonal ($a > b\sqrt{3}$) for all

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layered precursors, indicative of the systematic elongation of distorted Mn(III) octahedra along the *a* axis and of their ordered distribution. FTs of the EXAFS spectra consistently show similar Mn-Me_E distances in all layered precursors (Fig. S8). The systematic orthogonal symmetry of NiB layers is typical of octahedral layers containing few vacant layer sites (Drits et al., 1998; Lanson et al., 2000). Consistently, the presence of Ni(II) cations sorbed above or below such vacant sites could be excluded and at low Ni content Ni(II) cations are mostly sorbed at particle edges sharing either edges or corners with Mn octahedra (Simanova et al., 2015). Both XRD patterns and EXAFS spectra of layered precursors are modified with increasing Ni content, however. These changes reveal a significant decrease of the crystal size thus allowing more Ni to sorb at particle edges.

In addition, the increasing intensity of the \approx 4.6 Å diffraction maximum with increasing Ni content indicates that part of Ni initially introduced in solution actually precipitates as Ni (hydr)oxide. The additional presence of aqueous Ni in the interlayers of NiB15 and NiB20 is attested too by the presence of two planes of water molecules in their interlayers, consistent with its more negative hydration enthalpy compared to Na⁺ (Smith, 1977; Johnson and Post, 2006). XRD data clearly indicates that the relative proportions of these two Ni species weakly bound to layered precursors increase with increasing Ni content. All Ni species, including the two mentioned above, are readily mobilized by the acid treatment (Fig. 3c).

In a Ni-free system, a typical triclinic birnessite such as Bir converts readily and completely to an *a*-disordered todorokite (Wu et al., 2019). The sorption of low Ni(II) contents at birnessite particle edges, as in NiB5, does not hamper this birnessite-to-todorokite transformation. On the other hand, when increasing the Ni content, the presence of Ni(II), although weakly bound to the layered precursors to a large extent, significantly impacts their transformation during the reflux process (Figs. 3e, 5; Table 4). The first effect is an increased disorder of the tunnel structures, as evidenced by the decrease in intensity of the 761 cm⁻¹ band in the infrared spectra of reflux products (Fig. S9) and by the increasing proportion of non-ideal $3 \times n$ tunnel sizes forming trilling patterns (Fig. S10). In addition, platy crystals are visible in all reflux products, their relative abundance increasing with increasing Ni content with only minimal contributions of tunnel oxides being detected in NiT15 and NiT20 (Fig. 7b). In addition, analysis of Mn K-edge EXAFS (Figs. 5, 6 and S5; Table 4) shows that with increasing Ni content the proportion of phyllomanganates in reflux products increases at the expense of tectomanganates. Both the increase of the $I_{4.8}/I_{9.6}$ intensity ratio and the evolution of the layer-to-layer distance upon nitric acid treatment (Fig. 3e) indicate that asbolane, a phyllomanganate hosting an incomplete octahedral layer of metal (hydr)oxides in its interlayers, prevails in Ni-rich reflux products. These incomplete brucite-like octahedral layers are dissolved by the acid treatment, leading to birnessite-like structures exhibiting the observed evolution of their layer-to-layer distance (Fig. 3e). As reported in the early work of Chukhrov and coworkers on natural

asbolane (Chukhrov, 1982; Chukhrov et al., 1987), XRD diffraction patterns of this mineral are characterized by high $I_{4.8}/I_{9.6}$ intensity ratios. Such high ratios are commonly reported for "todorokite", especially when synthesized in the presence of metallic elements such as Co, Ni, Cu, Zn, Er, La, etc. (Shen et al., 1994; Tian et al., 1997; Ching et al., 1999; Luo et al., 1999). In the present study, the combined use of XRD simulation, anomalous XRD, HRTEM, thermogravimetric analysis, and nitric acid treatment allowed uncovering the actual nature of this seldom described phyllomanganate, but its formation may be far more common than previously reported in these experimental studies.

As shown in Fig. 2e, this $I_{4.8}/I_{9.6}$ intensity ratio strongly depends on the completeness of the interlayer brucite-like octahedral layer, and the ratio determined for NiT15 and NiT20 corresponds to a \approx 50% completeness of the Ni (OH)₂ octahedral layer. This interlayer occupancy is much higher than possibly achieved with the sole presence of Ni (II) in these octahedral layers as Ni/(Ni + Mn) ratios are 8–12% in these samples, with part of the Ni present as Ni (hydr)oxide. It is thus likely that Mg cations contribute to the electron density at the interlayer mid-plane, in addition to Ni. As Mg scattering factor is lower than that of Ni, the actual completeness of the interlayer brucite-like layer is likely higher than 50%.

4.2. Differentiation of asbolane and todorokite

Although the $I_{4.8}/I_{9.6}$ intensity ratio strongly depends on the completeness of the interlayer brucite-like octahedral layer, a 4.8 Å peak stronger than its 9.6 Å counterpart is a possible indication of the presence of the phyllomanganate asbolane, rather than the tectomanganate todorokite, in reaction products or in natural samples. The two compounds indeed share these two interplanar distances. With its two planes of interlayer H₂O molecules, buserite also exhibits similar periodicity but a gentle heating (to $\approx 100 \,^{\circ}$ C) should lead to the partial dehydration of interlayer cations and to the ≈ 7.2 Å periodicity typical of birnessite (Fig. S1 – Manceau et al., 2007b). Furthermore, buserite is unlikely to have a high enough content of interlayer cations to induce a $I_{4.8}/I_{9.6}$ ratio higher than 1.0.

Although values of this ratio higher than 1.0 are typically observed for asbolane, the partial exchange of Mg present in todorokite tunnels by cations having a higher scattering factor (such as Ni and other divalent metal cations) can also lead to such high $I_{4.8}/I_{9.6}$ ratios (Fig. 9), thus not allowing an unambiguous differentiation between asbolane and todorokite. In this search for an unquestionable differentiation between asbolane and todorokite, the nitric acid treatment proposed by Wu et al. (2019) thus appears as a simple and reliable method. In the case of asbolane, dissolution of the interlayer brucite-like layer induced by the treatment leads indeed to a collapse of the layer-to-layer distance to the ≈ 7.2 Å periodicity. On the contrary, periodicities are maintained for todorokite, the tectomanganate framework being unaltered, and positions of XRD maxima are thus unchanged.



Fig. 9. Intensity distributions calculated for ideal todorokite (ICDD #38-0475) with (a) only Mg in todorokite tunnels and (b) with Mg and Ni present in a 1:1 ratio in todorokite tunnels. In both cases, the model of Post et al. (2003) was modified assuming the complete filling of the O8 site by H₂O molecules (no cations) and the presence of 1.0 cation (Mg or Mg + Ni) in the Mg site.

4.3. The role of Mn(III) and interlayer species during the transformation

It is widely accepted that the presence of a high Mn(III) content (>25-30%) in layered precursors is required for their conversion to tectomanganates. This pivotal role is most likely related to the steric strains induced by the presence of Jahn-Teller distorted Mn(III) octahedra in the octahedral layers. Migration of these Mn(III) cations from layer to interlayer or layer kinking at these structurally weak points contribute to building tunnel walls (Grangeon et al., 2014, 2015; Atkins et al., 2014, 2016; Yang et al., 2018). In the present case, the ordered distribution of Mn (III) cations in rows parallel to the b axis and separated from each other along the *a* axis by two rows of Mn(IV)creates a structurally weak point favorable to the formation of tunnels walls (Atkins et al., 2014; Grangeon et al., 2014). Consistently, the initial Ni-free precursor Bir, a triclinic birnessite, fully converts to todorokite. Similar transformation of triclinic birnessite to tunnel structures was previously reported in the literature although 3×3 tunnels were not systematically obtained (Yang et al., 2018). Furthermore, the actual distribution and orientation of Mn(III) octahedra present in the MnO₂ layer do not appear to be key factors for the phyllomanganate-to-tectomanganate conversion as tectomanganates were obtained from hexagonal birnessite precursors in which Mn(III) octahedra are distributed and/or oriented at random (Grangeon et al., 2014, 2015; Atkins et al., 2014, 2016; Zhao et al., 2015). The absence of transformation of NiB precursors to tunnel structures thus does not appear to be related to their Mn (III) content, which is similar to or higher than that of the Ni-free Bir.

As layers of all NiB precursors are essentially devoid of Ni, both in the octahedral layer and sorbed above/below

vacant layer sites, the absence of transformation of these precursors to tunnel structures is thus likely due to kinetic effects favoring the formation of asbolane. Specifically, the hydrolysis and polymerization of hydrated Ni(II) as Ni(OH)₂ fragments in the interlayers of initial birnessite/ buserite (Fig. 7b; Table 5) appears to be kinetically favored over the build-up of tunnel walls which is essential to the formation of tectomanganates. In turn, the presence of interlayer Ni(OH)₂ fragments likely impedes the migration of Mn(III) from the layer to the interlayer thus preventing the conversion to tectomanganates. Such competing effects are supported also by the increase of (Mg + Ni)/Mn ratios with increasing Ni content (Table 1) and are likely to occur with a variety of other hydrolysable metal cations. Consistently, "todorokite" containing divalent metal cations and displaying high I_{4.8}/I_{9.6} ratios possibly indicate the presence of asbolane-like structure. These competing effects might contribute also to the prevalence of phyllomanganates over tectomanganates in natural Ni-rich environments (Wu et al., 2019).

4.4. Negative influence of Ni and Co on the phyllo-totectomanganate transformation: Contrasting mechanisms

Both in the present work investigating the possible transformation of Ni-containing layered precursors to tectomanganates and in the related study investigating a similar process for Co-containing layered precursors (Wu et al., 2019), tectomanganate formation was inhibited, essentially (Ni) or partially (Co) for Me-rich precursors. No negative effect is observed however for Me contents of $\approx 1-2$ wt.%, whatever their association with layered precursors. At such low Me content, Ni is either partially incorporated in the layer structure (Atkins et al., 2016) or sorbed at particle edges (this study), whereas Co is mainly structurally

incorporated in the octahedral layer (Wu et al., 2019). The origin of the negative impacts observed for high Me contents differs however for the two metals. In the present case, layered precursors contain sufficient amounts of Mn(III) (\approx 1/3) to trigger the transformation, as Ni(II) is not structurally incorporated in the precursors but rather present at particle edges (at low Ni content) and weakly bound, either as interlayer aqueous Ni(II) or as Ni (hydr)oxide, to Ni-rich precursors (NiB15 and NiB20). As discussed above, hydrolysis and polymerization of Ni(OH)₂ fragments in birnessite interlayers is kinetically favored over migration of Mn(III) from the layer to the interlayer, or layer kinking, leading to the formation of asbolane rather than tectomanganates.

In contrast and consistent with previous reports (Manceau et al., 1997; Yu et al., 2012; Kwon et al., 2013; Simanova and Peña, 2015; Yin et al., 2015), Co(II) is readily oxidized by birnessite during the formation of initial layered precursors with Co(III) being structurally incorporated in these precursors owing to the similar ionic radii of Co(III) and Mn(IV) (Shannon, 1976; Wu et al., 2019). The symmetry of layered precursors is thus increased from orthogonal to hexagonal with Co loading. As a consequence, the content of Mn(III) in Co-containing layered precursors is lowered and does not allow triggering the transformation to tunnel structures (Grangeon et al., 2014, 2015; Atkins et al., 2014, 2016). In both cases, little Me is released to solution during the reflux process of Me-rich layered precursors and part of Me cations are structurally incorporated in the reflux products ($\approx 20\%$ and $\approx 80\%$ for Ni and Co, from the nitric acid experiments and EXAFS simulation, respectively) thus enhancing their sequestration in Mn oxides.

4.5. The fate of Nickel during the transformation of layered precursors

Previous studies have consistently shown that Ni can be sequestered by birnessite, either within the octahedral layer or sorbed at vacant layers sites and/or at particle edges (Peacock and Sherman, 2007b; Grangeon et al., 2008, 2017; Peacock, 2009; Peña et al., 2010; Yin et al., 2012, 2014; Kwon et al., 2013; Simanova et al., 2015; Atkins et al., 2016; Lefkowitz and Elzinga, 2017), consistent with natural occurrences in Mn nodules (Bodeï et al., 2007; Manceau et al., 2007a, b; Peacock and Sherman, 2007a). In their literature review, Grangeon et al. (2017) showed that the fraction of Ni structurally incorporated in birnessite octahedral layers may be high (up to \approx 50%) only for low Ni content ($\approx 1\%$), as reported for natural samples (Bodeï et al., 2007; Manceau et al., 2007a, b; Peacock and Sherman, 2007a). Structural incorporation of Ni(II) in the octahedral layer appears to be enhanced by circumneutral pH conditions and time (Manceau et al., 2007b; Peacock and Sherman, 2007b; Peacock, 2009; Peña et al., 2010). Competitive sorption at TCS/DCS sites has been proposed also as a mechanism favoring structural incorporation in the octahedral layer (Lefkowitz and Elzinga, 2017). Some studies report the prevalence of Ni sorbed at TCS/DCS sites even for Ni/Mn ratios higher than 10%, however

(Manceau et al., 2007a, b; Peña et al., 2010; Yin et al., 2012; Simanova et al., 2015; Grangeon et al., 2017). In such cases, sorption of Ni(II) as inner sphere complexes is favored by the presence of vacant octahedral sites and is thus negatively correlated with the Mn(III) content. Such high Ni/Mn ratios may also be obtained with Mn(III)rich phyllomanganates through Ni sorption at particle edges, however (Simanova et al., 2015; Grangeon et al., 2017). Consistently, structural incorporation of Ni in vacancy-free NiB precursors appears very limited in the present study and Ni sorption at particle edges is limited to low Ni contents. In NiB layered precursors, Ni is rather present both as Ni (hydr)oxide and as hydrated cations in birnessite/buserite interlayers, as shown by the increased hydration of Ni-rich precursors with increasing Ni content (Figs. 1, 7a, and S1). The influence of Ni on phyllomanganate hydration has been reported also for natural samples in which buserite (or 10 Å vernadite) abundance increases with increasing Ni content (Bodeï et al., 2007; Manceau et al., 2007a).

As discussed above, hydrated Ni(II) initially present in birnessite interlayers are readily stabilized during the reflux process by forming Ni(OH)₂ fragments through hydrolysis (Fig. 7b; Table 5 – Lu et al., 2012), consistent with previous experiments with vacancy-free phyllomanganate precursors (Feng et al., 2001). Temperature stability of interlayer Ni $(OH)_2$ fragments (Fig. S3) is increased compared to that of primary hydrated species (Fig. S1). Polymerization of Ni(OH)₂ brucite-like octahedral layers in phyllomanganate interlayers is consistent also with the natural predominance of asbolane in natural Ni-rich environments (NiO \approx 10–20 wt.%, - Chukhrov, 1982; Chukhrov et al., 1987; Roque-Rosell et al., 2010; Ploquin et al., 2018). Although the Mn AOS of his synthetic samples (\approx 4.0) was favorable for Ni sorption at layer vacancy sites, Manceau et al. (2007b) also reported the overwhelming presence of Ni as Ni(OH)₂ at high Ni content (Ni/Mn ratio of ≈ 0.10 and ≈ 0.18 at pH 4 and 7, respectively), despite experimental conditions supposed to prevent Ni(OH)₂ precipitation. The reflux process appears to have little influence on the Ni (hydr)oxide initially formed together with the layered precursors, as evidenced by the stability of the corresponding maximum at ≈ 4.6 Å in the reflux products (Fig. 1b). Finally, part of Ni (≈ 20 %) is stabilized during the reflux process, as shown by the incomplete Ni release during acid treatment (Fig. 3d). Stabilized Ni is possibly structurally incorporated in the reflux products. On the contrary, Atkins et al. (2016) reported no structural incorporation of Ni(II) during the formation of todorokite from a Nibearing birnessite precursor containing ≈ 1 wt.% Ni present in similar proportions as sorbed and structurally incorporated species in hexagonal birnessite. They reported also a decrease in the Ni content of their reflux products, whereas in our experimental conditions stability of Ni/(Mn + Ni)ratios (Table 1) indicates that all Ni initially present in layered precursors is retained during the reflux process. Ni speciation in Ni-rich layered precursors thus appears to control the fate of this element during their subsequent transformation.

5. CONCLUSION

The present study reports on the association of weakly bound Ni species [hydrated Ni, Ni (hydr)oxides] with phyllomanganates, that possibly prevail in natural Ni-rich (>10% NiO) manganates. Whatever their Ni content, synthesized phyllomanganates systematically exhibit an orthogonal layer symmetry, indicative of a high Mn(III) content in octahedral layers and of a minimal number of octahedral layer vacancies. At high Ni(II) content, Ni is thus essentially present as charge-compensating hydrated Ni(II) and as Ni (hydr)oxide, whereas Ni(II) sorbed at particle edges prevail at low Ni(II) content. No Ni(II) structurally incorporated in the phyllomanganate structure was detected in the initial vacancy-free layered precursors.

The high Mn(III) content in Ni-free triclinic birnessite, a favorable condition for the conversion to tectomanganates, allows its complete transformation to todorokite upon reflux treatment. Absolane rather than tectomanganates forms during the reflux processing of Ni-rich phyllomanganates, however. Asbolane formation results from the kinetically favored polymerization of fragments of Ni(OH)₂ octahedral layers in birnessite interlayers. The structure of the initial manganate layers is essentially unaffected by this transformation.

Formation of asbolane, rather than todorokite, may actually be more frequent than reported in the literature owing to the similar periodicities in both compounds. High values (>1) of the $I_{4,8}/I_{9,6}$ intensity ratio may suggest the presence of this seldom described phyllomanganate. Acid nitric treatment, aiming at the dissolution of the islandlike interlayer Ni(OH)₂ layer, represents an easy and unambiguous way to differentiate asbolane from todorokite which is unaffected by the treatment.

The presence of both Co and Ni in synthesized phyllomanganates impedes their transformation to tectomanganates when refluxed, the origin of this similar impact being different for the two metals, however. Oxidation of Co(II) sorbed on the MnO₂ layer and the subsequent structural incorporation of Co(III) in the octahedral layer releases the steric strains related to the initial presence of Jahn-Teller distorted Mn(III) octahedra, which is essential for the phyllomanganate-to-tectomanganate conversion. In contrast, the presence of hydrated Ni(II) favors the polymerization of fragments of Ni(OH)2 octahedral layers in the phyllomanganate interlayers, that in turn prevents the formation of tectomanganates despite the favorable presence of Mn(III) in the octahedral MnO₂ layer. In both cases, "foreign" metal cations are retained in the refluxed products, with contrasting ratios of structural incorporation.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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RESEARCH DATA

Original data from the present study can be accessed at https://doi.org/10.17632/jznwn42xr6.2.

APPENDIX A. SUPPLEMENTARY MATERIAL

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Transformation of Ni-containing birnessite to tectomanganate:

Influence and fate of weakly bound Ni(II) species

Supplementary material

Geochimica Et Cosmochimica Acta

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Fig. S1. XRD patterns of fresh layered precursors after heating to 140 °C for different durations



Fig. S2. XRD patterns of reflux products (zoom on low-angle regions). Colors as in Fig. 1.



Fig. S3. XRD patterns of reflux products after heating to 140 °C for 12 hours. Colors as in Fig. 1.



Fig. S4. Intensity ratio of the intensity of 4.8 Å and 9.6 Å peaks ($I_{4.8}/I_{9.6}$) in reflux products as a function of Ni content and X-ray energy (see Fig. 2 for the data).



Fig. S5. Linear combination fitting of NiT Mn K-edge EXFAS spectra using corresponding NiB and Tod spectra as references for layered and tunnel structures, respectively.



Fig. S6 Simulations of Ni K-edge EXAFS spectra for layered precursors in both k- (a) and r-space (b).



Fig. S7. Interatomic distances calculated for a variety of Mn-Mn pairs and ideal todorokite structure (ICDD #38-0475). The Mn4-Mn4 distance (4.48 Å) which is not reported in the figure is similar to the Mn2-Mn2 distance. Figure modified from Fig. 1(b) in Wu et al. (2019).



Fig. S8. Fourier transforms (FTs) of Mn K-edge EXAFS showing modulus and imaginary parts (solid and dotted lines, respectively) in r-space for layered precursors.



Fig. S9. FTIR spectra of layered precursors and reflux products (top and bottom, respectively). Colors as in Fig. 1. Infrared spectra were collected for both layered precursors and reflux products using a Bruker VERTEX 70. For each sample, 64 spectra were collected and summed over the 4000–400 cm⁻¹ range with a 4 cm⁻¹ resolution (Zhao et al., 2015).



Fig. S10. HRTEM images of (a-b): Tod, (c-e): NiT1, (f-h): NiT5, (i-m): NiT1. Numbers in (b) and (d) indicate the number of MnO₆ octahedra defining tunnel width. Arrows in (i) and (l) indicate the direction of fringes bending. Micromorphology of the reflux products and lattice periodicities were observed a Jeol JEM2100F transmission electron microscope (TEM) operated at 200 kV.

Table S1

Coordination environment of Ni species in layered precursors. Ni-Me1 accounts for Ni-Ni pairs in NiO and for Ni-Mn_{DES}; Ni-Me2 accounts for Ni-Ni pairs in Ni(OH)₂ and for Ni-Mn_{DES} (probably). For this simulation, CN of Ni-O was set to 6 for all samples, whereas CNs of Ni-Me1 and Ni-Me2 were adjusted using a 0.5 step to optimize the fit to the data; all other parameters were refined.

	NiB5	NiB10	NiB15	NiB20
CN	6	6	6	6
R (Å)	2.04(0)	2.03(1)	2.04(0)	2.04(1)
σ^2 (Å ²)	0.005(0)	0.005(0)	0.005(0)	0.005(1)
CN	9	9	6	6
R (Å)	2.94(1)	2.92(2)	2.91(1)	2.94(2)
$\sigma^2({\rm \AA}^2)$	0.019(2)	0.018(3)	0.010(2)	0.015(4)
CN	6	6	5.5	4
R (Å)	3.08(2)	3.06(2)	3.06(1)	3.06(2)
$\sigma^2({\rm \AA}^2)$	0.013(2)	0.013(2)	0.014(1)	0.010(3)
	-5.4(7)	-6.0(9)	-5.1(8)	-4.7(1.0)
	0.01	0.01	0.01	0.02
	CN R (Å) σ^{2} (Å ²) CN R (Å) σ^{2} (Å ²) CN R (Å) σ^{2} (Å ²)	NiB5 CN 6 R (Å) 2.04(0) σ² (Ų) 0.005(0) CN 9 R (Å) 2.94(1) σ² (Ų) 0.019(2) CN 6 R (Å) 3.08(2) σ² (Ų) 0.013(2) σ² (Ų) 0.013(2)	NiB5 NiB10 CN 6 6 R (Å) 2.04(0) 2.03(1) σ² (Ų) 0.005(0) 0.005(0) CN 9 9 R (Å) 2.94(1) 2.92(2) σ² (Ų) 0.019(2) 0.018(3) CN 6 6 R (Å) 3.08(2) 3.06(2) σ² (Ų) 0.013(2) 0.013(2) σ² (Ų) 0.013(2) 6.0(9)	NiB5 NiB10 NiB15 CN 6 6 R (Å) 2.04(0) 2.03(1) 2.04(0) σ² (Ų) 0.005(0) 0.005(0) 0.005(0) CN 9 9 6 R (Å) 2.94(1) 2.92(2) 2.91(1) σ² (Ų) 0.019(2) 0.018(3) 0.010(2) CN 6 5.5 1.01 σ² (Ų) 0.013(2) 3.06(1) 3.06(1) σ² (Ų) 0.013(2) 0.013(2) 0.014(1) σ² (Ų) 0.013(2) 0.014(1) 1.01

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