



Transformation of the phylломanganate vernadite to tectomanganates with small tunnel sizes: Favorable geochemical conditions and fate of associated Co

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Abstract

The present work uncovers the geochemical control on the nature (tunnel size) of the tectomanganates formed from layered precursors, and thus provides insights into the formation of Mn oxides in natural environments. Large tunnel sizes are favored under circum-neutral conditions, whereas low pH conditions favor the formation of tectomanganates with smaller tunnel sizes. Both the increased proportions of Mn(III) in vernadite/birnessite layers resulting from low pH conditions and the subsequent enhancement of Mn(III) disproportionation during subsequent transformation contribute to the formation of tectomanganates with smaller tunnel sizes. The fate of foreign elements during the phylломanganate-to-tectomanganate mineral transformation is another important aspect of this mineral transformation, together with the impact of these elements on the transformation. Layered and tunnel Mn oxides have indeed a pivotal influence on the geochemical cycling of transition metals, including Co, that possess a strong affinity for these mineral species. The present experimental work shows that the formation of todorokite (3 × 3 tunnel size), hollandite (2 × 2), or nsutite (intergrown 1 × 1 and 1 × 2 fragments) is essentially unaffected by limited Co-enrichment (≤5 at.%) of the initial phylломanganate structure. Higher Co contents reduce the content of Jahn-Teller distorted Mn(III) octahedra in layered precursor and hamper the phylломanganate-to-tectomanganate transformation. Finally, Co is retained in the structure of todorokite and hollandite during their formation under circum-neutral conditions whereas part (~20%) of the Co present in layered precursors is expelled out of the framework and/or sorbed to nsutite formed under acidic conditions. This effect is induced by the reduced stability of Co(III) octahedra when the relative proportion of corner-sharing linkages increases. In turn, this effect influences Co structural incorporation in different Mn oxides and its potential release to solution.

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1. INTRODUCTION

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Manganese oxides are common in ferro-manganese nodules or polymetallic crusts found in aquatic and terrestrial

environments (Chukhrov et al., 1981, 1987; Hein et al., 1988; Bodeř et al., 2007). Among these oxides, layered oxides (phylломanganates: birnessite, vernadite, busserite, asbolane, and lithiophorite) and tectomanganates with medium [2×2 : hollandite (Ba^{2+}) and cryptomelane (K^+)] and large tunnels [3×3 : todorokite] have attracted special attention. The sustained interest for these minerals results from their ubiquity in a variety of natural environments, their common enrichment in transition metals and rare-earth elements (Burns and Burns, 1978; Ostwald, 1984; Hein et al., 1988; Post, 1999), and from their key influence on the transport and fate of organic and inorganic contaminants (McKenzie, 1989; Post, 1999).

By contrast, tectomanganates with smaller tunnel sizes (1×1 : pyrolusite; 1×2 : ramsdellite; intergrowth of 1×1 and 1×2 fragments: nsutite – Zwicker et al., 1962; Post, 1999) are abundant in manganese ore deposits worldwide (Zwicker et al., 1962; Turner and Buseck, 1983; Parc et al., 1989; Nicholson, 1992 and references therein). In addition to the widespread use of manganese in metallic alloys, these economic deposits are mined for the electrochemical performance of the tectomanganates with smaller tunnel sizes (Turner and Buseck, 1983). They usually result from intense long-term supergene weathering of terrestrial and marine deposits (see for example the review of Nicholson, 1992 and references therein), and more especially of Mn-bearing carbonates and silicates under humid tropical or subtropical climates (Mücke et al., 1999; Dekonink et al., 2019). In these contexts, tectomanganate formation is not meant to imply layered precursors, despite contradictory observations in lateritic profiles (Parc et al., 1989). Formation of tectomanganates such as nsutite from layered precursors was also reported experimentally (Tu et al., 1994; Luo et al., 2018). More generally, tectomanganate formation from phylломanganate precursors has attracted special attention (Golden et al., 1986, 1987; Feng et al., 2004; Shen et al., 2005; Bodeř et al., 2007), owing to the major contribution of this formation pathway to the natural abundance of tectomanganates. Mn(III) migration out of the octahedral MnO_2 layers to release steric strains arising from the Jahn-Teller distortion of Mn(III) octahedra and/or layer kinking at these structurally weak points appear as the most likely mechanisms for the formation of tectomanganates having large and medium tunnel sizes, such as todorokite (Atkins et al., 2014) and hollandite/cryptomelane (Grangeon et al., 2014, 2015).

In nodules and crusts, Mn oxide enrichment in transition metals and rare earth elements occurs through sorption, isomorphous substitution (structural incorporation), or intimate association with (oxyhydr)oxides, and resulting concentrations of these elements may reach several percents (Taylor, 1968; Burns and Burns, 1978; Manceau et al., 1987, 2007; Hein et al., 1988; Post, 1999; Marcus et al., 2004; Bodeř et al., 2007; Peacock and Sherman, 2007; Asavin et al., 2015). Together with Ni, Co is among the most common transition metals found in association with, and structurally incorporated in, both layered and tunnel Mn oxides (Taylor, 1968; Manceau et al., 1987, 1997; Hein et al., 1988; McKenzie, 1989; Kay et al., 2001;

Asavin et al., 2015; Simanova and Peña, 2015; Wu et al., 2019). (Co,Ni) enrichment is for example typical for the so-called “hydrogenetic” Mn oxides (Bonatti et al., 1972; Choi and Hariya, 1992; Nicholson, 1992). Specifically, surface-catalyzed oxidation of Co(II) by Mn(III) and subsequent migration of Co(III) in former Mn(III) octahedral sites account for Co enrichment in phylломanganates (Manceau et al., 1997; Simanova and Peña, 2015). Co(III)-for-Mn(III) substitutions are sterically favored and release steric strains arising from the presence of Jahn-Teller distorted Mn(III) octahedra, and thus negatively influence birnessite conversion to todorokite (Drits et al., 1997; Manceau et al., 1997; Gaillot et al., 2007; Yin et al., 2015; Wu et al., 2019). In addition, high Co concentrations (>5 at.%) in layered Mn oxides commonly leads to the formation of asbolane as described by Chukhrov et al. (1987), precluding further the transformation to tunnel structures. When this transformation occurs, that is experimentally for Co content < 7.7 at.%, Co is essentially retained in the solid phase, being mainly present in octahedral sites sharing only edges with adjacent octahedra (Mn1/Mn3 sites of todorokite – Wu et al., 2019; Wegorzewski et al., 2020). Co may also be structurally incorporated in hollandite/cryptomelane (Li et al., 2015). Although favored in tectomanganates with large and medium tunnel sizes, Co contents of 1–2 at.% [Co/(Co + Mn)] have also been reported in natural pyrolusite (e.g., Lambiv Dzemua et al., 2013).

The fate of Co during the transformation from layered precursors to tectomanganates with medium-to-small tunnel sizes remains poorly documented, however. The present work thus aimed at preparing nsutite and hollandite from the same Co-containing vernadite precursor, a phylломanganate common in soils (Chukhrov et al., 1981; McKenzie, 1989; Vodyanitskii, 2006, 2009), sediments (Bodeř et al., 2007), and deep-sea Mn nodules (Chukhrov et al., 1987; Hein et al., 1988), using a reflux treatment mimicking natural processes (Feng et al., 2004; Atkins et al., 2014). Layered precursors and reflux products were characterized using chemical analysis, X-ray diffraction, Mn/Co K-edge X-ray absorption spectroscopy, and nitric acid treatment. Transformation processes were determined through the characterization of intermediate reaction products. In addition, special attention was paid to decipher the fate of Co during the formation of the two tectomanganates. The present work thus provides significant new insight into the geochemical conditions enhancing the formation of tectomanganates with contrasting tunnel sizes, and the crystal-chemical features driving the fate of associated foreign metal elements in natural environments.

2. MATERIALS AND METHODS

2.1. Preparation of Co-containing vernadite

Vernadite was synthesized according to a modified version of the protocol reported by Villalobos et al. (2003). Specifically, 300 mL of a 0.2 M KMnO_4 solution, which was boiled and cooled to room temperature, were dropped into 300 mL of a 0.667 M NaOH solution at a 5 mL/min rate. Next, 300 mL of a solution containing 23.7 g of

MnCl₂·4H₂O and 2.37 g of CoCl₂·6H₂O [Co/(Co + Mn) = 5.3 at.%) were added to the mix at a 5 mL/min rate. The resulting suspension was left to settle for 2 hrs (final pH of the supernatant was 4.32), filtered and washed with ultrapure water (18.2 MΩ·cm) until its conductivity was < 20 μS/cm. Part (~110 mL) of this suspension was freeze dried, this sample being referred to as CoV.

2.2. Transformation of Co-containing vernadite to nsutite

Half of the remaining CoV suspension was sampled, and its volume adjusted to 500 mL with ultrapure water. This suspension was then equilibrated at pH 4 for 3 days using HNO₃ or NaOH. A 100 mL aliquot was sampled and washed, the resulting sample being referred to as CoV-pH4. Most of the remaining 400 mL suspension was transferred to a flask and refluxed at 100 °C for 2 days, during which 30 mL aliquots were sampled at different time intervals. After cooling to room temperature, the final pH of the supernatant was 5.52. The samples were named CoV-pH4-Re-*nn*, *nn* being the reflux time. For example, the final product is referred to as CoV-pH4-Re-2d. In addition, sample CoV-0.5 M H⁺-Re was obtained by refluxing an aliquot of CoV in a 0.5 M HNO₃ aqueous solution.

2.3. Transformation of Co-containing vernadite to hollandite

The other half of the remaining suspension was contacted with 500 mL of a 1 M NaCl solution for 1 day (Zhao et al., 2015), and washed until conductivity was < 20 μS/cm. The resulting homoionic suspension was then resuspended in a 0.5 M BaCl₂ solution for 1 day and washed again with ultra pure water until conductivity was < 20 μS/cm. The volume of this suspension was adjusted to 500 mL, and its pH equilibrated to 7.50 ± 0.05 for 1 day. The resulting sample is referred to as CoV-Ba. 400 mL of the CoV-Ba suspension was then refluxed at 100 °C for 2 days, leading to sample CoV-Ba-Re. After cooling to room temperature, the final pH of the supernatant was 6.12. All the above solid samples were retrieved through centrifugation and freeze-dried.

2.4. Sample characterization

X-ray diffraction was performed on a Bruker D8 diffractometer, equipped with Cu Kα radiation (λ = 1.5418 Å) and a SolXE solid-state detector (Baltic Scientific Equipments). Data was collected from 2–90 °2θ Cu Kα in step mode with 40 s counting times per 0.04 °2θ step. Sample chemical composition was measured in duplicate using inductively-coupled plasma spectrometry (ICP-OES Varian 720ES) after dissolution of the samples in NH₂OH.HCl.

To assess the relative proportion of elements structurally incorporated in the solids and sorbed at their surface, 0.1 g aliquots were acid-washed in 250 mL of a 1 M HNO₃ solution under moderate stirring at room temperature (23 °C). 5 mL aliquots of this suspension were sampled at different time intervals and immediately filtered through a 0.22 μm membrane to determine the release of Mn, Co, and Ba, the contents of which were analyzed by ICP-OES.

Mn/Co K-edge Extended X-ray Absorption Fine Structure (EXAFS) spectra were collected at room temperature on the 1W1B beamline of the Beijing Synchrotron Radiation Facility (BSRF) to determine Mn/Co local environments in both layered precursors and reflux products. Data was collected in transmission (Mn K-edge) or fluorescence (Co K-edge) mode, a metallic Mn/Co foil being used as reference for energy calibration. Athena and Artemis softwares were used to process and simulate EXAFS spectra (Ravel and Newville, 2005). Path simulations for Mn K-edge EXAFS were performed over the 1–4 Å (R + ΔR) range in r-space and over 3.1–12.7 Å⁻¹ in k-space, with an amplitude reduction factor (*S*₀²) of 0.8 (Grangeon et al., 2010). Path simulations for Co K-edge EXAFS were performed over the 1–3.8 Å (R + ΔR) range in r-space and over 3.2–10.5 Å⁻¹ in k-space, with an amplitude reduction factor (*S*₀²) of 0.877 (Yin et al., 2014).

The Mn average oxidation state (AOS) of layered precursors and reflux products was determined by applying a linear combination fitting method to the Mn K-edge XANES 1st derivative data, that is specifically designed for the determination of Mn valence states in mixed-valent Mn oxides (the so-called Combo method – Manceau et al., 2012). The reference compounds used for Mn(II), Mn(III) and Mn(IV) were those of the original study, the uncertainty on the relative proportion of the different Mn oxidation states being ± 4% (Manceau et al., 2012).

3. RESULTS

3.1. Elemental composition

Co/(Co + Mn) ratios are about the same for all analyzed samples at 5.05–5.28 at.% (Table 1). Limited Co, or Mn, is thus removed, exchanged, or desorbed from the solid phase during pH equilibration, cation exchange, or reflux. On the other hand, all Na⁺ initially present in CoV is replaced by protons following equilibration at pH 4, when only ~1/3 of K initially present is replaced. An additional ~1/5 of K is released during the reflux of CoV-pH4. For CoV-Ba, after two-steps (Na⁺ and Ba²⁺) of exchange, no K and only minor Na were detected. The significant increase in Mn content from layered precursors to their reflux products is most likely related to their contrasting H₂O contents and the related “dilution” effect. Birnessite/vernadite contains indeed ~2% adsorbed H₂O and ~7% interlayer H₂O (Lanson et al., 2002) compared to ~3.0% of adsorbed and lattice H₂O in nsutite (Said, 2020).

3.2. Mn average oxidation state

During the reflux process, the Mn AOS of Co-containing layered precursors increases systematically (Table 2). This increase is enhanced under low pH conditions as shown by the respective AOS determined for CoV-0.5 M H⁺-Re-2d and CoV-pH4-Re-2d (3.90 and 3.72, respectively). This increase contrasts with the Mn AOS stability reported during birnessite conversion to

Table 1
Elemental composition of selected samples.

Sample	Mn wt.%	Co wt.%	Co at.%	Ba at.%	K at.%	Na at.%
CoV	56.1(7)	3.24(8)	5.11	n.d.	4.7	1.4
CoV-pH4	55.7(4)	3.23(6)	5.12	n.d.	3.0	n.d.
CoV-pH4-Re-1h	60.9(4)	3.55(6)	5.16	n.d.	2.8	n.d.
CoV-pH4-Re-2d	65.1(2)	3.71(2)	5.05	n.d.	1.6	n.d.
CoV-Ba	48.4(9)	2.89(6)	5.28	8.5	n.d.	3.5
CoV-Ba-Re	56.3(0)	3.36(0)	5.28	7.9	n.d.	2.6

Note: at.% values are calculated as $Me/(Co + Mn)$; n.d. stands for not detected.

Table 2

Relative proportions of Mn (II), (III), and (IV) valence states determined by fitting Mn K-edge XANES spectra 1st derivative (Combo method – Manceau et al., 2012) and resulting Mn average oxidation state (AOS).

Sample	Mn(II) at.%	Mn(III) at.%	Mn(IV) at.%	Mn-AOS
CoV	9	30	60	3.51
CoV-pH4	10	26	65	3.55
CoV-pH4-Re-1h	6	22	72	3.66
CoV-pH4-Re-3h	5	25	70	3.65
CoV-pH4-Re-2d	2	23	75	3.72
CoV-0.5 M H ⁺ -Re-2d	1	8	91	3.90
CoV-Ba	6	24	70	3.63
CoV-Ba-Re	1	18	80	3.79

todorokite under similar reflux conditions (Atkins et al., 2014; Wu et al., 2019).

3.3. Powder XRD patterns

All three layered precursors (CoV, CoV-pH4, and CoV-Ba), display a weak and broad peak at $\sim 7 \text{ \AA}$ (Fig. 1a), indicative of a limited periodicity of MnO₆ octahedral layer stacking along the c* axis (Drits et al., 1997; Villalobos et al., 2006; Grangeon et al., 2008; Lanson et al., 2008; Yin et al., 2012). XRD patterns obtained for these three samples display also an asymmetric peak at $\sim 2.43 \text{ \AA}$ and

a broad maximum at $\sim 1.41 \text{ \AA}$, whose position and shape are typical of octahedral layers having hexagonal symmetry. Together, these three peaks are typical for vernadite (Fig. 1a – Chukhrov et al., 1987; Drits et al., 2007). In addition, XRD pattern of CoV-pH4 shows weak modulations at 3.95, 2.14, and 1.64 \AA , indicative of a minor nsutite contribution (ICDD#17-0510) resulting from low-pH conditions maintained for several days (Fig. S1).

Compared to those of their layered precursors, XRD patterns of the reflux products display additional reflections (Fig. 1b). For example, the XRD pattern of CoV-pH4-Re-2d display reflections at 4.02, 2.45, 2.15, and 1.67 \AA ,

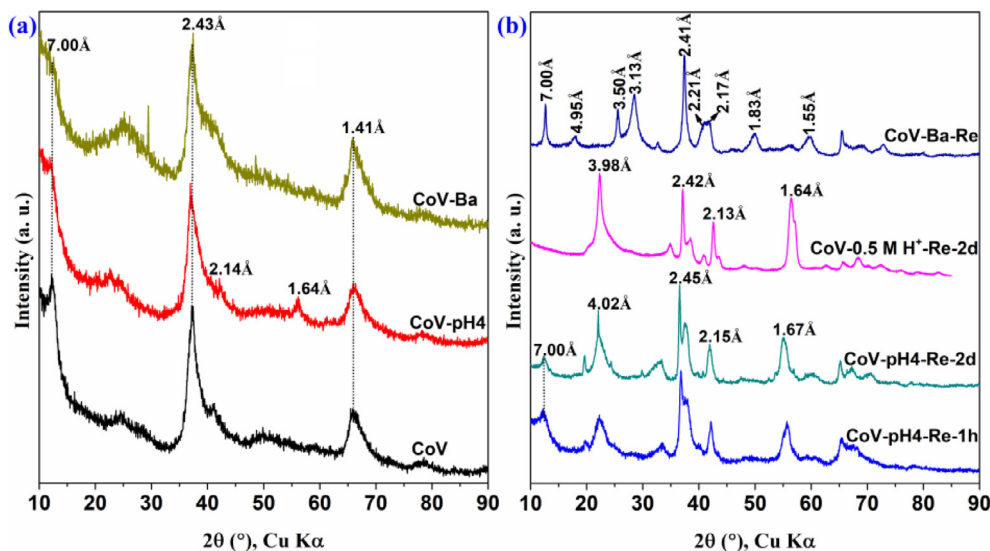


Fig. 1. XRD patterns of layered precursors (a) and of their reflux products (b).

consistent with the prominent lines of manganian nsutite (ICDD#14-0614 – Fig. S1). Nsutite consists in the intergrowth of pyrolusite and ramsdellite fragments, thus displaying both 1×1 and 1×2 tunnel sizes (Zwicker et al., 1962; Turner and Buseck, 1983; Post, 1999), its manganian equivalent containing presumably a significant amount of Mn(II,III) (Zwicker et al., 1962). XRD pattern of CoV-pH4-Re-1h is essentially similar to that of CoV-pH4-Re-2d, except for a lower signal-to-noise ratio (Fig. 1b). Mineralogical transformation from the vernadite precursor to manganian nsutite thus occurs very early during the reflux process, the initial crystallization stage being followed by a steady crystal growth. In addition, the persistence of the ~ 7.0 Å reflection in XRD patterns of both CoV-pH4-Re-1h and CoV-pH4-Re-2d indicates the presence of residual vernadite.

Although the XRD pattern of CoV-0.5 M H^+ -Re-2d looks like that of the former, reflections of CoV-0.5 M H^+ -Re-2d are shifted towards higher angles, compared to those of the latter. Peak positions in CoV-0.5 M H^+ -Re-

2d are consistent with the presence of ideal nsutite (ICDD#17-0510), rather than manganian nsutite, consistent with the higher Mn AOS determined in this sample, compared to CoV-pH4-Re-2d. Finally, XRD pattern of CoV-Ba-Re is similar to that of hollandite (ICDD #38-0476 – Fig. S1), a tectomanganate with a uniform 2×2 tunnel size (Post et al., 1982; Miura, 1986).

3.4. Mn local environment

Mn K-edge EXAFS spectra collected for both layered precursors and reflux products (Fig. 2a) have similar frequency, except over the 7.0 – 9.5 Å⁻¹ “indicator region” (Marcus et al., 2004). Specifically, layered precursors have troughs at ~ 7.5 Å⁻¹ while their reflux products display weak peaks at the same frequency. On the contrary, layered precursors display sharp peaks of similar intensities at ~ 8.2 Å⁻¹, whereas reflux products display much weaker peaks at this frequency, that of CoV-0.5 M H^+ -Re-2d almost vanishing. Mn K-edge spectra of the three layered

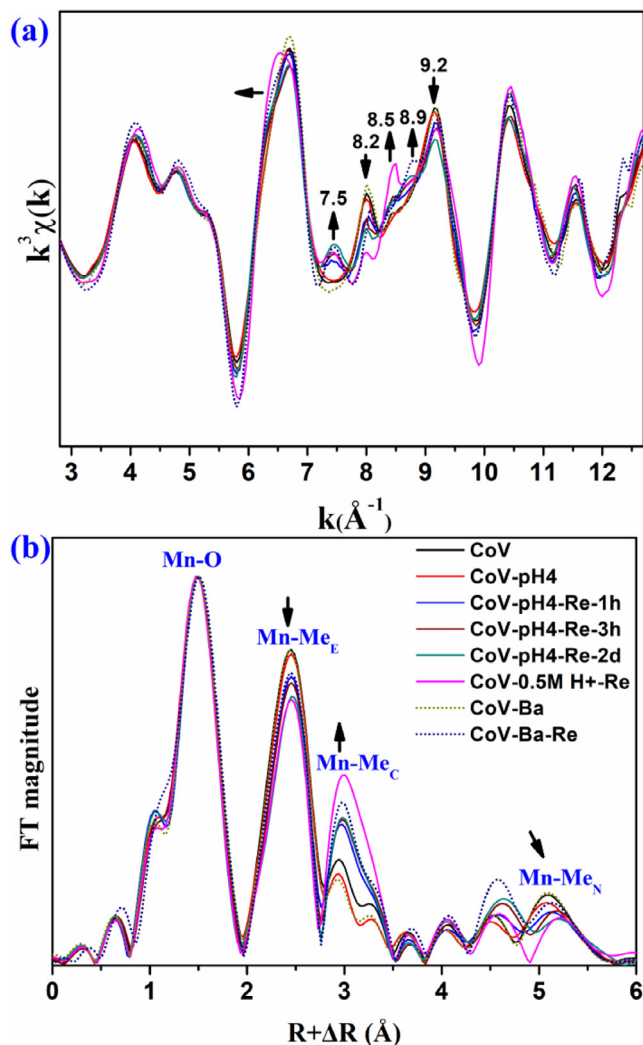


Fig. 2. Mn K-edge EXAFS spectra (a) and corresponding Fourier transforms (FT – b). FT intensities are normalized to the ~ 1.5 Å ($R + \Delta R$) maximum that corresponds to the first Mn-O shell.

precursors are essentially similar over the 8.5–9.2 Å⁻¹ range, whereas spectra of CoV-0.5 M H⁺-Re-2d and CoV-Ba-Re display distinct peaks at ~8.6 Å⁻¹ and ~8.9 Å⁻¹, respectively (Fig. 2a). The contrasting profiles of Mn K-edge spectra over this “indicator region” are indicative of different contents and distributions of Mn (III) and Mn(IV) in layered precursors, and of variable tunnel sizes in reflux products (McKeown and Post, 2001; Bargar et al., 2005; Webb et al., 2005; Toner et al., 2006; Bodei et al., 2007).

The Fourier transforms (FTs) of these EXAFS spectra (Fig. 2b) display three main peaks at ~1.5, ~2.5, and ~3.0 Å (R + ΔR) that correspond to the nearest Mn-O, edge-sharing Mn-Me (Me = Mn, Co), and corner-sharing Mn-Me shells, respectively (McKeown and Post, 2001; Webb et al., 2005). Positions of these three peaks are similar for all samples but the relative intensities of the latter two peaks vary between layered precursors and reflux products (Fig. 2b). The intensity at ~2.5 Å (R + ΔR) decreases indeed from layered precursors to reflux products whereas an opposite trend is visible at ~3.0 Å (R + ΔR). Both trends are maximized for CoV-0.5 M H⁺-Re-2d, that displays lowest intensity at ~2.5 Å (R + ΔR) and highest at ~3.0 Å (R + ΔR), indicative of low numbers of edge-sharing linkages (ESL) and of high numbers of corner-sharing linkages (CSL). Peaks at ~5.2 Å (R + ΔR) represent the second-neighbor Mn-Me in octahedral sheets (Bargar et al., 2005; Webb et al., 2005; Toner et al., 2006; Wu et al., 2019). Consistent with the trend observed for the first edge-sharing Mn-Me shell, this peak decreases in intensity from layered precursors to reflux products, indicative of the decreasing proportion and/or size of layered fragments.

Coordination numbers (CN) and distances typical for the different shells are derived from the simulation of Mn K-edge EXAFS spectra and of their FTs in both k and r spaces (Table S1 – Grangeon et al., 2010). In layered precursors, Mn atom have ~4 nearest Mn neighbors, a number much lower than the theoretical value of 6 even taking experimental uncertainties into account. Small particle size, and related edge effect (Bargar et al., 2005), and presence of vacant octahedral sites likely account for the observed reduction of CN (Grangeon et al., 2010). On the other hand, CN of Mn-Me CSL increases from layered precursors to their reflux products leading to a remarkable increase in the CN_C/CN_E ratio, consistent with a phyllo-manganate-to-tectomanganate conversion (Kim et al., 2003; Webb et al., 2005; Wu et al., 2019). There is no significant evolution of the interatomic distances for the different shells between layered precursors and reflux products.

3.5. Co local environment

Co K-edge EXAFS spectra of selected samples display similar frequencies but contrasting amplitudes over the k-range probed (Fig. 3a), amplitudes decreasing in the order vernadite (CoV) to hollandite (CoV-Ba-Re), and to nsutite (CoV-pH4-Re-2d). The FTs of these Co K-edge EXAFS spectra (Fig. 3b) display three maxima at ~1.5, ~2.5, and

~3.0 Å (R + ΔR) that correspond to Co-O first shell, and Co-Me ESL and CSL, respectively (Simanova and Peña, 2015; Wu et al., 2019). FTs of CoV and CoV-pH4-Re-2d are essentially alike, except for a stronger intensity at ~2.5 Å (R + ΔR) in the former sample. The maximum at ~2.5 Å (R + ΔR) is weakest for CoV-Ba-Re whereas the intensity at ~3.0 Å (R + ΔR) is highest, thus indicating different numbers of ESL and CSL, consistent with the CN_C/CN_E ratios derived from the simulation of Co K-edge EXAFS spectra (Fig. S3 and Table S2). Low Co-Me CSLs indicate the minor presence of Co sorbed at vacancy sites (Co^V) in CoV, whereas most Co (~81%) are structurally incorporated species (Co^S), consistent with previous reports (Manceau et al., 1997; Kwon et al., 2013; Simanova and Pena, 2015; Wu et al., 2019).

3.6. Nitric acid treatment

Consistent with previous reports (Wu et al., 2019, 2020), XRD patterns of CoV, CoV-pH4-Re-2d, and CoV-Ba collected after the nitric acid treatment display minor evolutions relative to the original ones (Fig. 4a), indicative of the limited modification of Mn oxide structure. Weak peaks at 4.9, 3.5, 3.1, 2.2, 1.8, and 1.55 Å, consistent with those of hollandite (ICDD #38-0476), are observed in CoV XRD pattern, however, thus confirming the formation of tunnel structures during the acid-wash treatment. Hollandite reflections are also detected in CoV-Ba XRD pattern after the acid treatment, although weaker than in CoV pattern. Reflection positions and intensities are essentially unchanged in CoV-Ba-Re pattern after the nitric acid treatment. By contrast, although the overall XRD profile of CoV-pH4-Re-2d appears unaffected by the acid treatment, reflections are systematically shifted towards smaller distances by this treatment. This minor contraction of the unit cell is consistent with the transformation from manganooxide to ideal nsutite and is most likely related to the release to solution of Mn(II,III) from the solid (Zwicker et al., 1962).

Nitric acid treatment is an effective method to distinguish metal species in Mn oxides, as nitric acid is able to extract exchangeable and adsorbed metal cations without damaging significantly (limited reduction essentially) the manganese oxide framework (Wu et al., 2019). In the present work, Mn (Fig. 4b) and Co (Fig. 4c) releases do not reach a maximum after one week of acid treatment, although release rates are much reduced compared to the initial rate. Mn is preferentially released from layered precursors compared to their reflux products, suggesting lower contents of soluble Mn(II,III) in the latter, consistent with results of the Combo method. Mn release, and Mn release rate after 24 hrs, is maximum for CoV.

During the acid-wash treatment, CoV-pH4-Re-2d releases ~49% of its Co content to solution, more than its layered precursor (~37%), and far more than CoV-Ba-Ex (~31%) and CoV-Ba-Ex-Re (~20%). The initial fast release of Co, that can be quantified from the intersect on the y-axis of the linear fitting to Co release during the final stages (>=24 hrs) of treatment, will be hereafter considered as sorbed Co (Co^V), whereas the remaining Co will be

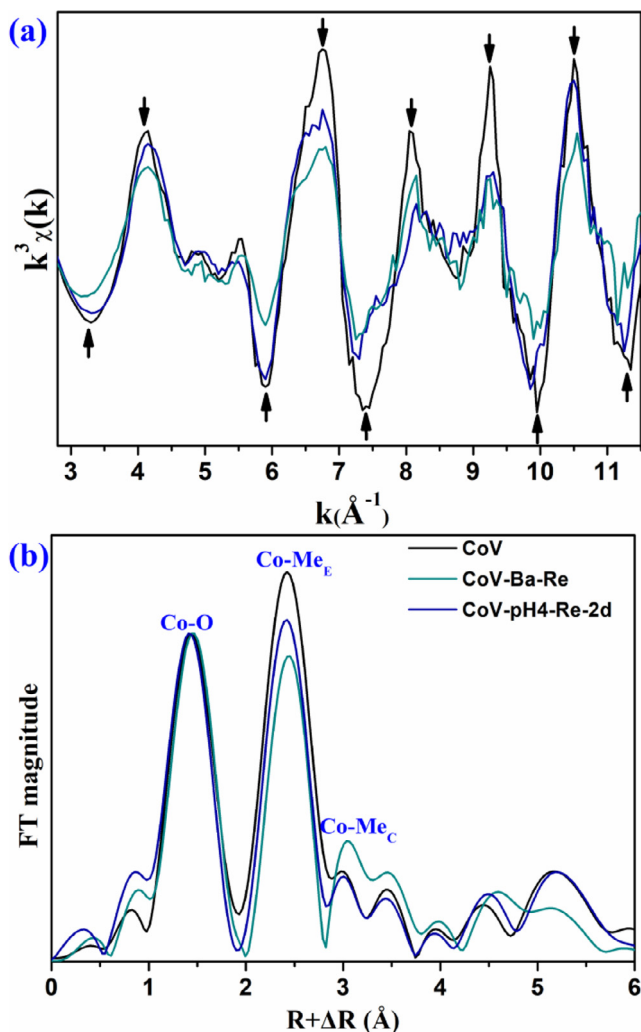


Fig. 3. Co K-edge EXAFS spectra (a) and corresponding Fourier transforms (FT – b). Arrows in (a) highlight the amplitude decrease from CoV to CoV-Ba-Re and to CoV-pH4-Re-2d. FT intensities are normalized to the $\sim 1.5 \text{ \AA}$ ($R + \Delta R$) maximum that corresponds to the first Co-O shell.

considered as structurally incorporated Co (Co^{S}). Accordingly, Co releases as a function of time indicate that Co^{V} and Co^{S} account for ~ 20 and $\sim 80\%$, respectively, in layered precursors (Table 3), consistent with the results derived from fitting the Co K-edge EXAFS data and with previous reports of Co-speciation in birnessite and vernadite (Kwon et al., 2013; Simanova and Peña, 2015; Yin et al., 2015; Wu et al., 2019). On the other hand, Co^{V} and Co^{S} account for ~ 60 and $\sim 40\%$ of the Co species in CoV-pH4-Re-2d, although its overall Co-content is similar to that of layered precursors (Table 1). The contrasting distributions of Co species observed in CoV and CoV-pH4-Re-2d indicate that Co speciation is significantly altered along the mineralogical transformation from vernadite to manganoan nsutite. Specifically, part of Co ($\sim 20\%$) is likely released from vernadite layers and sorbed to reaction product. On the contrary, the similarity of CoV and CoV-Ba-Re Co K-edge EXAFS spectra suggests that relative proportions of Co^{V} and Co^{S} were not modified significantly during conversion of Ba-exchanged vernadite (CoV-Ba) to hollandite

(CoV-Ba-Re – Table 3), consistent with previous report on todorokite formation from Co-containing layered precursors (Wu et al., 2019). The contrasting contents of Co^{V} and Co^{S} in CoV-Ba-Re and CoV-pH4-Re-2d are not correlated to their respective contents of CSL and ESL (Table S2). The high Co release ratio of CoV-pH4-Re-2d associated to its low $\text{CN}_{\text{C}}/\text{CN}_{\text{E}}$ ratio possibly results from a high content of double-corner sharing species.

Ba initially present as a hydrated exchangeable Ba^{2+} in CoV-Ba interlayers is readily released during the acid treatment (Fig. 4d). On the other hand, only $\sim 8\%$ of Ba is released from CoV-Ba-Re after a similar treatment, consistent with the coordination of Ba with O atoms forming hollandite 2×2 tunnels (Post et al., 1982; Miura, 1986). This behavior is specific to large Ba^{2+} cations (1.35 \AA ionic radius in sixfold coordination – Shannon, 1976) when present in 2×2 tunnels as smaller cations, such as Cd^{2+} (0.78 \AA ionic radius in sixfold coordination), could be present in tunnel sites as hydrated cations (Randall et al., 1998).

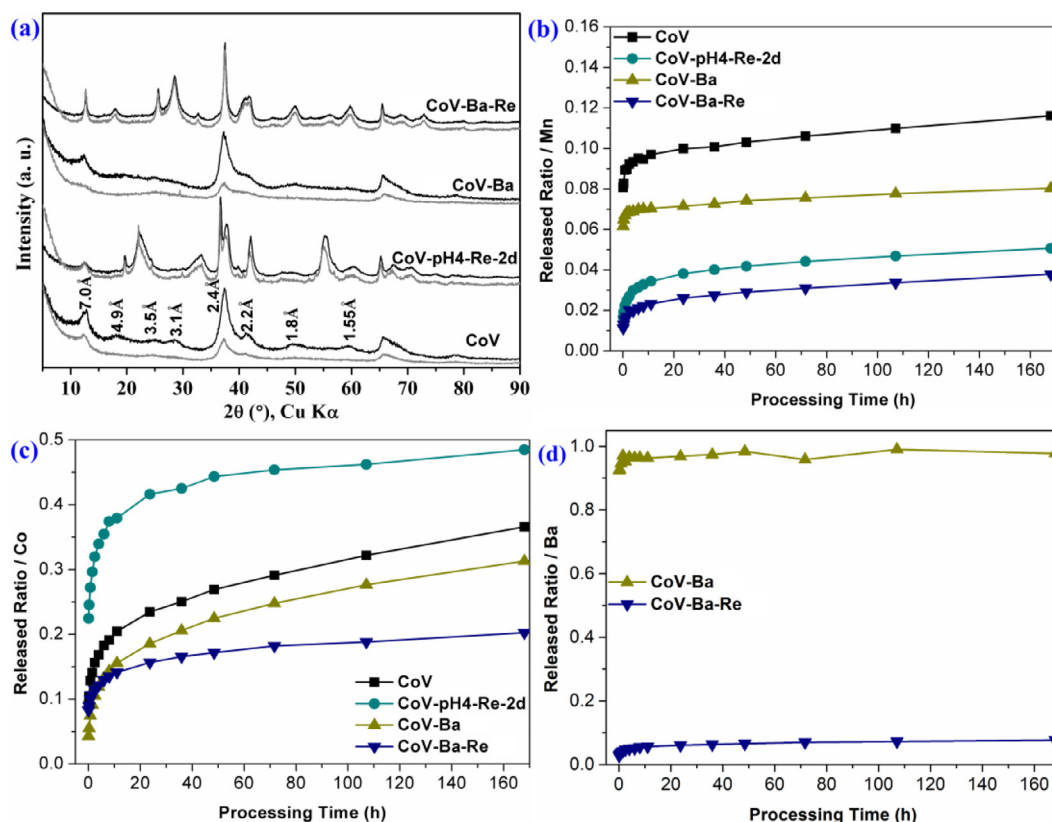


Fig. 4. Influence of the nitric acid-wash treatment on layered precursors and reflux products. XRD patterns before (gray) and after (black) the acid treatment (a). Elemental release from the solid during the treatment: Mn (b); Co (c); Ba (d). Co release curves are divided in two segments: data points before (not included) and after (included) 24 hrs define the fast and slow release stages, respectively. The slow release section is linearly fitted, the intercept of the equation approximating the amount of adsorbed species (CoV: $\%Co_{rel} = 0.0009x + 0.2211$, $R^2 = 0.9845$; CoV-pH4-Re-2d: $\%Co_{rel} = 0.0004x + 0.4138$, $R^2 = 0.9254$; CoV-Ba: $\%Co_{rel} = 0.0009x + 0.1775$, $R^2 = 0.9646$; CoV-Ba-Re $\%Co_{rel} = 0.0003x + 0.1552$, $R^2 = 0.9459$).

Table 3
Relative proportions of Co species in layered precursors and reflux products.

Samples	Co ^S /Co at. %	Co ^V /Co at. %	Co ^S /(Mn + Co) at. %	Co ^V /(Mn + Co) at. %
CoV	78	22	3.99	1.12
CoV-pH4-Re-2d	59	41	2.98	2.07
CoV-Ba	82	18	4.33	0.95
CoV-Ba-Re	84	16	4.44	0.84

*Note: The proportions of Co^V are determined from the intercept of the linear fit to the slow release portion of the elemental release curves during the nitric acid treatment (Fig. 4).

4. DISCUSSION

4.1. Formation of tectomanganates with different tunnel sizes from vernadite precursors

In the present work, both nsutite (intergrowth of 1×1 and 1×2 tunnels) and hollandite (2×2 tunnel size) were obtained from the reflux of Co-containing vernadite. The formation of tectomanganates with tunnel sizes smaller than 2×2 complements previous reports of both hollandite/cryptomelane and todorokite (3×3 tunnel size) formation from layered precursors and supports the hypothesis of the major role of vernadite as a precursor

in the formation of natural tectomanganates (Burns and Burns, 1978; Bodeř et al., 2007).

Environmental pH appears to strongly affect the nature of tectomanganates formed through its influence on tunnel dimension, as shown by the formation of pyrolusite, romanechite, and OMS-5 (1×1 , 2×3 , and 2×4 tunnel sizes, respectively) from the same layered precursor by varying pH conditions from 1, to 7, and to 13, respectively (Shen et al., 2005). Consistently, todorokite is obtained experimentally under neutral and weakly alkali or acidic conditions (pH 4.8–9 – Cui et al., 2005; Atkins et al., 2014), and formation of a tectomanganate with 4×4 tunnel size was reported at pH 6–8 from a layered vernadite

precursor in the presence of Mn(II) (Yang et al., 2018). To the authors' knowledge, there is no evidence of todorokite formation under low pH conditions ($\text{pH} < 4.8$), and Grangeon and coworkers showed that hollandite/cryptomelane formation was favored by the initial equilibration of the layered precursors at low pH (Grangeon et al., 2014, 2015), consistent with previous reports (Tu et al., 1994; Zhang et al., 2011). In the present work, hollandite was also formed at room temperature during the acid treatment (1 M H^+) of the vernadite precursor as well as under neutral condition during the reflux process (initial and final pH values: 7.50 and 6.12, respectively). In this case, Ba^{2+} or K^+ template the formation of 2×2 tunnels of hollandite and cryptomelane, respectively, owing to their appropriate radii (Post et al., 1982; Miura, 1986).

The influence of pH on tectomanganate tunnel size may be understood from the mechanisms driving the phylломanganate-to-tectomanganate reaction. The presence of Mn(III) in phylломanganate octahedral layers induces indeed steric strains owing to the Jahn-Teller distortion of Mn(III)O_6 octahedra (Drits et al., 1997; Gaillot et al., 2007). Mn(III) octahedra thus represent weak structural points, especially when aligned to minimize steric strains (Gaillot et al., 2007). Independent of the actual mechanism leading to the formation of tunnel walls (layer kinking or Mn(III) migration to the interlayer – Atkins et al., 2014, 2016; Grangeon et al., 2014, 2015), an increased proportion of Mn(III) favors the transformation process. Lowering the equilibration pH of the layered precursor thus appears to have a positive effect on the formation of tectomanganate with small tunnel sizes owing to the induced increase of Mn(III) in vernadite layers (Grangeon et al., 2014) and to the subsequent enhancement of Mn(III) disproportionation reaction (Manceau et al., 1997; Lanson et al., 2000). In the present case, disproportionation appears as an important step of the transformation as shown by the decreased content of Mn(III) in reflux products compared to Mn(IV) (Table 2). Enhancement of Mn(III) disproportionation and migration of Mn(II,III) from the layer to solution at low pH is supported by the presence of ideal and manganoan nsutite in CoV-0.5 M H^+ -Re and CoV-pH4-Re-2d, respectively. Mn(II,III) relative proportion is indeed significantly lower in ideal nsutite than in manganoan nsutite (Zwicker et al., 1962), and in hollandite and todorokite (Wu et al., 2019). Similarly, the nitric acid treatment of CoV-pH4-Re-2d releases Mn(II,III) to solution and induces the conversion of manganoan nsutite to its ideal form (Fig. 4a).

Temperature may also influence the tunnel size in formed tectomanganates. Hollandite is formed indeed during CoV acid treatment (1 M H^+) at ambient temperature, whereas manganoan nsutite is formed during the reflux treatment of CoV-pH4 at 100°C . Consistent with this hypothesis, formation of cryptomelane was reported after aging at room temperature of vernadite equilibrated at pH 3 (Grangeon et al., 2014). The enhanced formation of tectomanganates with increasing temperature is possibly related to the increased rate of Mn(III) disproportionation (Hanf et al., 2020) and release of Mn(II,III) to solution. Low-temperature formation of tectomanganates is likely

common in natural environments, however, especially in anoxic and suboxic conditions owing to the presence of Mn(II,III) species (Yang et al., 2018).

4.2. Influence of Co on tectomanganate formation, fate of Co during this process, and Co speciation in newly formed tectomanganates

It is commonly accepted that Co is mainly ($\sim 80\%$ – 100%) present as incorporated species in layered Mn oxides (Manceau et al., 1997; Kwon et al., 2013; Yin et al., 2014, 2015; Simanova and Peñna, 2015; Wu et al., 2019). Co (III) usually replaces Mn(III) isomorphously and its presence thus decreases the steric strains induced by the presence of these Jahn-Teller distorted octahedra in the layer (Manceau et al., 1997; Gaillot et al., 2007; Li et al., 2014; Yin et al., 2015) and impairs in turn the phylломanganate-to-tectomanganate transformation (Wu et al., 2019). Accordingly, Co contents higher than ~ 5 at. % in birnessite precursors were recently shown to hamper their transformation to todorokite and to lead to increasingly disordered $3 \times n$ ($3 < n$) products (Wu et al., 2019). For $\text{Me}/(\text{Me} + \text{Mn})$ ratios higher than 20 at. % ($\text{Me} = \text{Co}, \text{Ni}$), Co forms, often in association with Ni, an octahedral sheet sandwiched in between MnO_2 layers leading to the minerals asbolane or lithiophorite, depending on the Al content (Chukrov et al., 1987; Manceau et al., 1987; Llorca and Monchoux, 1991; Roqué-Rosell et al., 2010; Lambiv Dzemua et al., 2013; Ploquin et al., 2019). The presence of this octahedral sheet has also been shown to impede the transformation of layered Mn oxides to tectomanganates (Wu et al., 2020).

For Co contents in layered precursors similar to those in the present study [~ 0.05 Co/(Co + Mn) molar ratios], tunnel structures are systematically formed, however, and the overall Co content of the precursors is maintained in the reflux products, whatever their nature (Wu et al., 2019). Evolution of Co local environment differs however depending on the mineralogy of the reflux product (todorokite, hollandite, nsutite). In todorokite, Co exhibits essentially ESL, being mostly located in Mn1/Mn3 sites (Wu et al., 2019). In hollandite, the proportion of structural Co is similar to that of its layered precursor (CoV-Ba and CoV-Ba-Re in Table 3), although the CN_C/CN_E ratio is significantly increased in the reflux product (from 0.36 in CoV to 0.96 in CoV-Ba-Re – Table S2). Although to a lower extent, this ratio is increased also in nsutite (0.50 in CoV-pH4-Re-2d – Table S2), that also contains $\sim 20\%$ more Co sorbed to the solid phase, as shown by the nitric acid treatment (~ 20 and $\sim 40\%$ Co^V in CoV and CoV-pH4-Re-2d, respectively – Table 3).

The contrasting behavior of Co in the reflux products most likely results from their structure, and more especially from the relative proportions of ESL and CSL. Tectomanganates with different tunnel sizes display indeed contrasting CN_C/CN_E ratios. In ideal todorokite, Mn2/Mn4 sites display 4 CSL and 4 ESL (Fig. S5), whereas Mn1/Mn3 sites only have 6 ESL, the ratio between the two sets of sites being constant at 2 (McKeown and Post, 2001). In hollandite, there is only one type of sites, similar to todorokite

Mn2/Mn4 sites and thus displaying 4 CSL and 4 ESL. Although less favorable than sites displaying mostly ESL, these sites are also suitable to host structural Co. Finally, nsutite possesses two types of octahedral sites, one similar to the Mn2/Mn4 sites of todorokite (4 CSL and 4 ESL) whereas the other site displays 8 CSL and 2 ESL (Fig. S5), as in pyrolusite (McKeown and Post, 2001). Depending on the relative proportions of pyrolusite and of ramsdellite fragments in nsutite, the ratio of these two sites varies significantly.

As tunnel size decreases from todorokite to nsutite, the CN_C/CN_E ratio thus increases significantly. The predominant occurrence of Co in Mn1/Mn3 sites of todorokite (Wu et al., 2019) is consistent with the increased stability of Co^S compared to Co^V in phylломanganates shown by DFT calculations (Kwon et al., 2013). Although less favorable than sites displaying mostly ESL, hollandite Mn sites and possibly equivalent todorokite Mn2/Mn4 sites, are also suitable to host structural Co, as shown by the similar proportion of structural Co in CoV-Ba and CoV-Ba-Re (Table 3). On the contrary, octahedral sites in nsutite pyrolusite-like fragments are most likely not appropriate to accommodate Co, owing to their higher CN_C/CN_E ratio. As a result, ~20% of Co initially present in CoV-pH4 migrate out of the solid during the transformation process, before being sorbed on the newly formed tectomanganate. Natural Co contents in tectomanganates having small tunnel sizes are thus lower than those found in tectomanganates with medium-to-large tunnel sizes (e.g., Taylor et al., 1964; Lambiv Dzemua et al., 2013).

As a consequence, Co is preferentially enriched in natural phylломanganates compared to tectomanganates (Taylor et al., 1964; Taylor and McKenzie, 1966; Taylor, 1968; Burns and Burns, 1977; Chukhrov and Gorshkov, 1980; Chukhrov et al., 1980; Chukhrov, 1982; Ostwald, 1984; Llorca, 1987; Manceau et al., 1987, 1992; Buret and Vanbrabant, 2015). When possible, the subsequent

transformation of Co-rich phylломanganates will lead to structures exhibiting large tunnel sizes, such as a-disordered todorokite, in which Co can remain structurally incorporated. Transformation operating at lower pH values will result in formation of tunnel structures having smaller tunnel sizes, however. In this case, part of the Co initially present in the phylломanganate will be released from the octahedral layers and/or sorbed at the surface of newly formed tectomanganates, thus potentially favoring subsequent release of Co to solution.

5. CONCLUSION

Depending on environmental conditions, phylломanganate transformation may lead to tectomanganates exhibiting a variety of tunnel sizes (Fig. 5), even at low temperature, and the present study provides significant new insights into the geochemical conditions controlling tunnel size of formed tectomanganates. Large tunnel sizes are favored under circum-neutral conditions (pH 4.8–9), the actual tunnel size being in this case influenced by the nature of interlayer cation in the layered precursor (Mg: todorokite, K/Ba: cryptomelane/hollandite). Under acidic conditions (pH < 4.8), both the increase of Mn(III) content in the layers of phylломanganate precursors, and the subsequent enhancement of Mn(III) disproportionation during their transformation contribute to the formation of tectomanganates with small tunnel sizes.

In turn, the fate of Co structurally incorporated in layered precursors is also influenced by the pH conditions of the transformation. For Co contents lower than 5–10 at. %, the transformation to tectomanganates is essentially unaffected and most Co remains structurally incorporated in the newly formed tectomanganates formed under circum-neutral conditions (todorokite, hollandite, cryptomelane). Under acidic conditions, part of Co present in layered precursors is released to solution and/or sorbed at

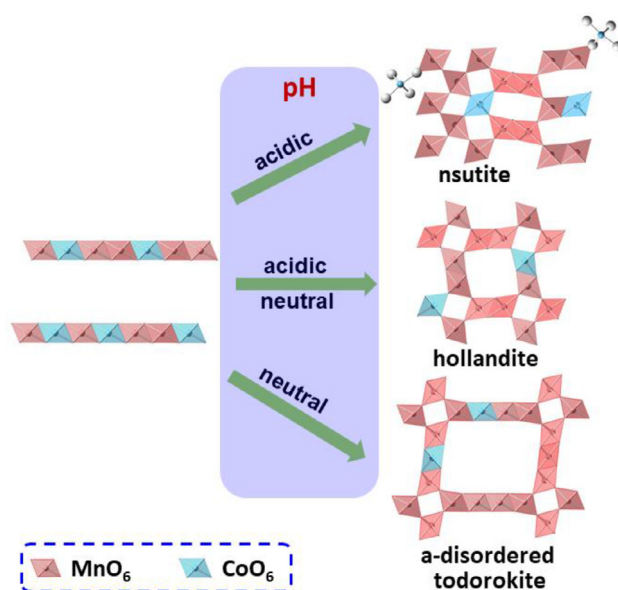


Fig. 5. Schematic of Co-containing vernadite transformation to tectomanganates with various tunnel sizes and behaviours of Co.

the surface of the tectomanganates having small tunnel sizes (nsutite, pyrolusite) owing to the reduced stability of Co(III) octahedra when the relative proportion of corner-sharing linkages increases. As Co(III) replaces Mn(III) isomorphously, its presence decreases the steric strains induced by the presence within the octahedral layer of these Jahn-Teller distorted octahedra and impairs the phyllomanganate-to-tectomanganate transformation when its content exceeds 5–10 at.% (Wu et al., 2019). The present results thus account for the natural prevalence of Co in phyllomanganates compared to tectomanganates, and in tectomanganates having large tunnel sizes compared to small ones, in which Co occurs as sorbed species to a large extent. By unraveling Co affinity for different Mn oxides, the present study thus contributes to a better understanding of the cycling of Co, and more generally of transition metals associated to environmentally relevant Mn oxides.

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

Research Data associated with this article contains: Chemical compositions; XAS data at Mn and Co K-edges; XRD data; Chemical analyses upon nitric acid treatment. Dataset can be accessed at <https://doi.org/10.17632/27369v78sv.2>.

APPENDIX A. SUPPLEMENTARY MATERIAL

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.gca.2020.12.021>.

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Supporting information to

Transformation of the phylломanganate vernadite to tectomanganates with small tunnel sizes: Favorable geochemical conditions and fate of associated Co

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This PDF file includes:

Supplementary Figures S1 – S5

Supplementary Tables S1 – S2

References

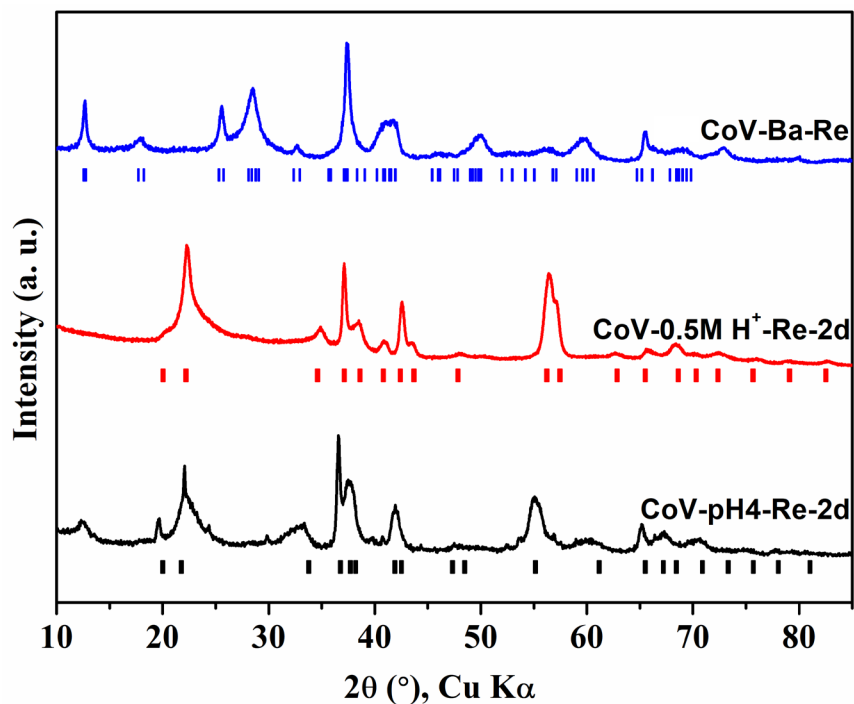


Fig. S1. X-ray diffraction patterns of the reflux products. Blue, red and black ticks indicate the position of reflections for hollandite (ICDD #38-0476), nsutite: (ICDD #17-0510), and manganian nsutite (ICDD #14-0614, not replaced), respectively.

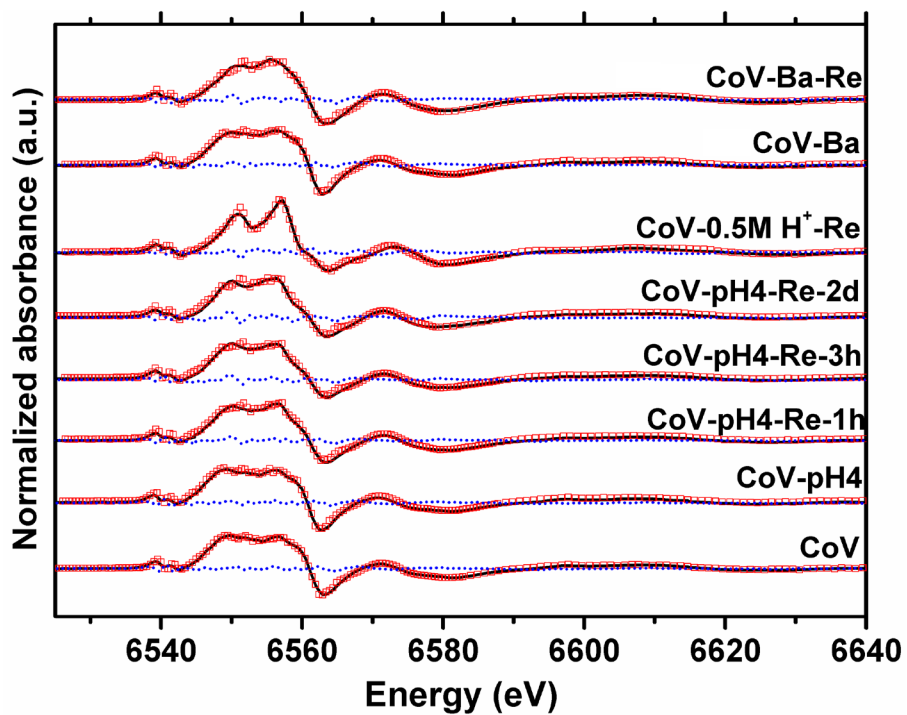


Fig. S2. First derivative of Mn K-edge XANES spectra (solid lines) of layered products and their reflux products, their best fits (open squares) obtained with the Combo method (Manceau et al., 2012) and difference plots (dotted lines).

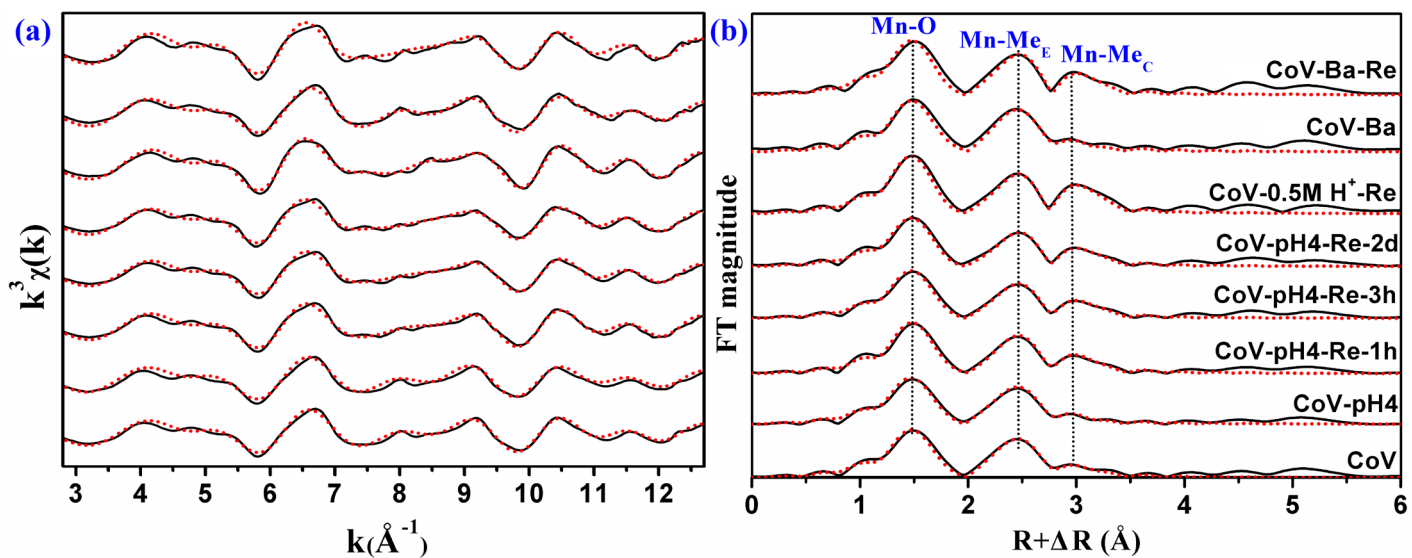


Fig. S3. (a) Mn K-edge EXAFS spectra (solid black lines) and their best fits (dotted red lines) and (b) corresponding Fourier Transforms (FTs – solid black lines) and their best fits (dotted red lines) for layered products and their reflux products

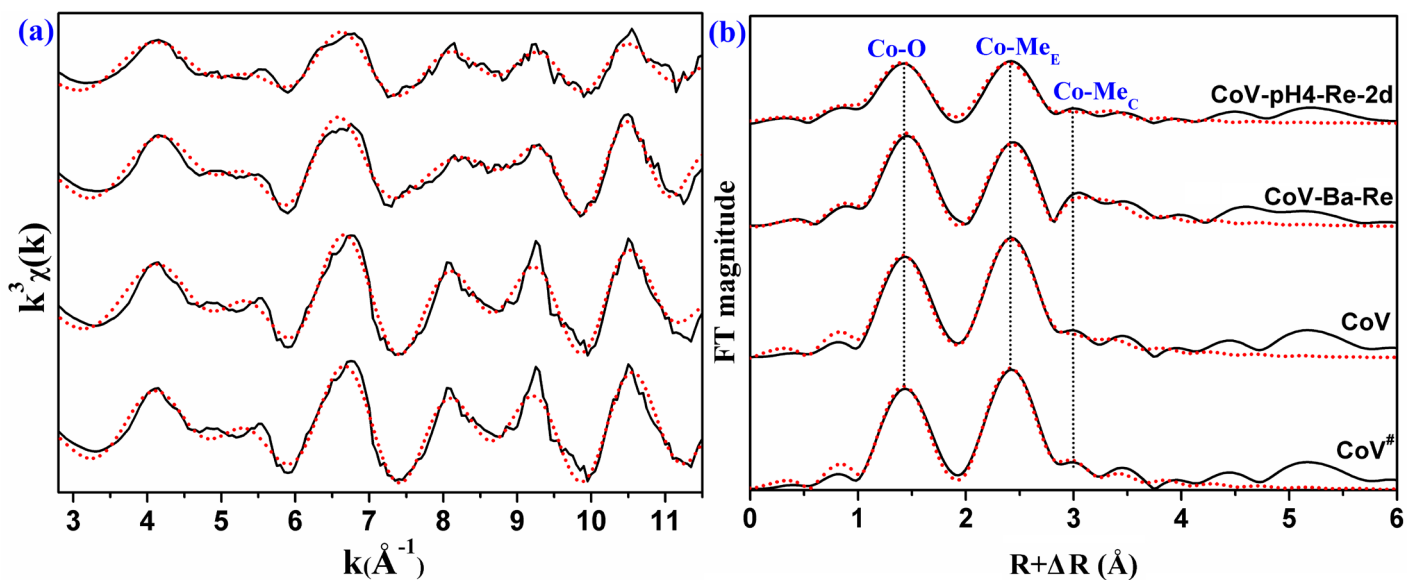


Fig. S4. (a) Co K-edge EXAFS spectra (solid black lines) and their best fits (dotted red lines) and (b) corresponding Fourier Transforms (FTs – solid black lines) and their best fits (dotted red lines) for layered products and their reflux products

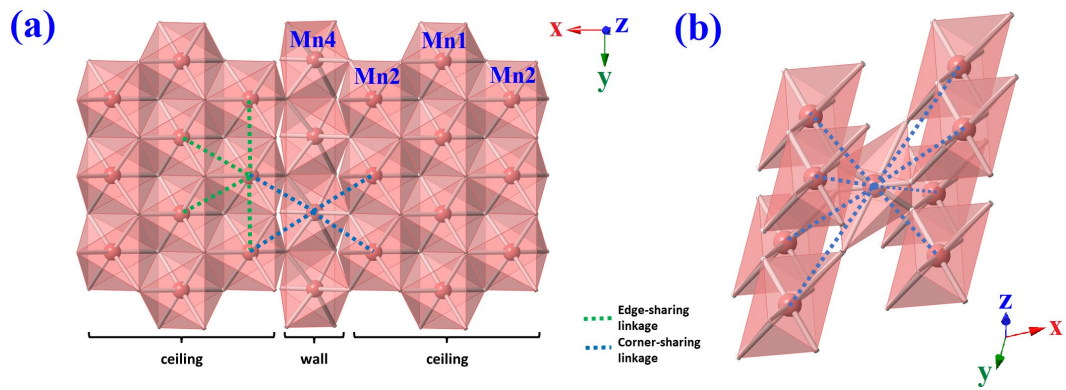


Fig. S5. Mn sites configurations in todorokite crystal structure (a) and that in pyrolusite (b). (a) is modified from [Wu et al. \(2019\)](#)

Table S1

Structural parameters derived from Mn K-edge EXAFS fitting over 1-3.8 Å (R+ΔR) in r-space

Atomic pairs		CoV	CoV-pH4	CoV-pH4-Re- 1h	CoV-pH4-Re- 3h	CoV-pH4-Re- 2d	CoV-0.5MH ⁺ -Re- 2d	CoV-Ba	CoV-Ba-Re
Mn-O _{1st}	CN	4.5(5)	4.6(5)	4.5(5)	4.6(37)	4.7(32)	5.3(5)	4.7(53)	5.43(9)
	R (Å)	1.90(1)	1.90(4)	1.90(4)	1.90(1)	1.90(0)	1.90(0)	1.90(0)	1.91(1)
	σ ² (Å ²)	0.004(1)	0.004(1)	0.003(1)	0.004(1)	0.004(1)	0.004(1)	0.004(1)	0.003(1)
Mn-O _{2nd}	R (Å) ^{a)}	3.63(3)	3.61(4)	3.60(4)	3.62(3)	3.61(3)	3.62(3)	3.60(4)	3.62(56)
Mn-Me _E	CN	4.0(6)	4.1(6)	3.3(5)	3.3(5)	3.2(4)	3.5(5)	4.1(7)	4.2(95)
	R (Å)	2.88(1)	2.88(2)	2.88(1)	2.88(1)	2.89(1)	2.88(1)	2.88(1)	2.89(1)
	σ ² (Å ²)	0.006(1)	0.006(1)	0.005(1)	0.006(1)	0.006(1)	0.005(1)	0.006(1)	0.007(2)
Mn-Me _C	CN	2.8(6)	2.7(6)	3.5(7)	3.7(6)	3.8(6)	5.1(7)	2.6(7)	4.9(1.4)
	R (Å)	3.48(3)	3.47(2)	3.46(1)	3.46(1)	3.46(1)	3.45(1)	3.46(3)	3.47(2)
CN _C /CN _E		0.71	0.66	1.05	1.09	1.19	1.44	0.64	1.16
ΔE ₀ (eV)		-5.4(1.1)	-5.9(1.1)	-4.8(1.1)	-5.2(1.1)	-5.2(1.0)	-5.6(1.2)	-4.8(1.0)	-4.5(1.5)
R factor (%)		0.02	0.01	0.02	0.02	0.01	0.01	0.02	0.05

Note: Coordination numbers (CN) and Debye-Waller factors (σ²) of the two Mn-O shells were refined jointly, as were Debye-Waller factors of the two Mn-Me shells.

Table S2

Structural parameters derived from Co K-edge EXAFS fitting over 1-3.8 Å (R+ΔR) in r-space.

Atomic pairs		CoV	CoV-Ba-Re	CoV-pH4-Re-2d	CoV [#]
Co-O _{1st}	CN	6	6	6	5.2(1.0)
	R (Å)	1.90(1)	1.91(4)	1.91(1)	1.90(1)
	σ ² (Å ²)	0.004(1)	0.005(1)	0.008(1)	0.002(1)
Co-O _{2nd}	R (Å) ^{a)}	3.67(6)	3.62(4)	3.64(7)	3.45(9)
Co-Me _E	CN	6.3(1.2)	3.7(1.0)	3.4(5)	6*f
	R (Å)	2.83(1)	2.86(1)	2.83(1)	2.83(1)
	σ ² (Å ²)	0.005(1)	0.004(1)	0.005(1)	0.003(2)
Co-Me _C	CN	2.3(1.2)	3.6(1.0)	1.7(1.0)	6*(1-f)
	R (Å)	3.49(4)	3.45(2)	3.46(4)	3.33(10)
CN _C /CN _E		0.36	0.96	0.50	f=0.81(13)
ΔE ₀ (eV)		-4.0(2.3)	-1.2(2.1)	-4.8(2.6)	-2.8(1.7)
R factor (%)		0.01	0.02	0.03	0.01

Note: Coordination numbers of Co-O_{1st} and Co-O_{2nd} were set to 6, and a parameter f representing the proportion of structurally incorporated Co species was introduced during the simulation of CoV[#]

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