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Nucleation and growth of feitknechtite from nanocrystalline vernadite precursor

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Abstract: Vernadite is a nanocrystalline manganese oxide, which controls the fate of many trace elements in soils and sediments through sorption and oxidative-degradation mechanisms. This exceptional reactivity directly results from its crystal structure, which may however evolve upon contact with redox-sensitive species. Understanding these changes is a prerequisite to predict and model the geochemical cycle of trace elements in the environment. Here, the structural and morphological modifications affecting synthetic nanocrystalline vernadite (δ -MnO₂) upon contact with increasing concentrations of Mn²⁺ were investigated using wet chemistry, synchrotron X-ray diffraction and transmission electron microscopy. Fresh δ -MnO₂ crystals had an Mn oxidation state of 3.94 ± 0.05 and a ~ 10 Å layer-to-layer distance. Crystal size was ~ 10 nm in the layer plane, and ~ 1 nm perpendicular to that. Upon contact with aqueous Mn²⁺ under anoxic conditions, δ -MnO₂ crystals underwent several morphological and mineral evolutions, starting with the stacking, perpendicular to the layer plane, of δ -MnO₂ crystals to form crystals ~ 10 nm \times 2 nm which were then subjected to oriented aggregation both along and perpendicular to the layer plane to form lath-like crystals with dimensions of ~ 100 nm \times 20 nm. Finally, these laths stacked perpendicular to the layer plane to form synthetic feitknechtite (β -MnOOH) crystals with sizes up to ~ 100 nm \times 500 nm when the Mn²⁺ loading reached 31.9 mmol g⁻¹. Structural transformation from δ -MnO₂ to synthetic feitknechtite was detected at Mn²⁺ loading equal to or higher than 3.27 mmol g⁻¹.

These mechanisms are likely to influence the geochemical fate of trace elements in natural settings where Mn²⁺ is abundant. Firstly, the systematic increase in crystal size with increasing Mn²⁺ loading may impact the sorption capacity of vernadite and feitknechtite by reducing the density of reactive edge sites. Secondly, the fate of trace elements initially sorbed at the vernadite surface is unclear, as they could either be released in solution or incorporated into the feitknechtite lattice.

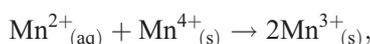
Key-words: vernadite; δ -MnO₂; feitknechtite; β -MnOOH; manganese oxide; X-ray diffraction; transmission electron microscopy; nucleation; sorption; crystal growth.

1. Introduction

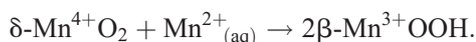
Manganese (Mn) oxides are ubiquitous in surficial environments, where they play a key role in the geochemical cycles of major and trace elements, as they are highly reactive towards nutrients (e.g., Xin *et al.*, 2016), metals (e.g., Halbach, 1986; Bellanca *et al.*, 1996; Manceau *et al.*, 2004, 2007, 2014; Marcus *et al.*, 2004; Boonfueng *et al.*, 2006; Fuller & Bargar, 2014), actinides (e.g., Kunzendorf & Friedrich, 1976; Koppi *et al.*, 1996; Duff *et al.*, 1999) and organic matter and organic molecules (e.g., Cheney *et al.*, 1996; Nasser *et al.*, 2000; Barrett & McBride, 2005; Chang Chien *et al.*, 2009; Johnson *et al.*, 2015). Vernadite, the nanocrystalline and

turbostratic analogue to birnessite (Giovanoli, 1980), is one of most abundant (Taylor *et al.*, 1964; Taylor, 1968; Ross *et al.*, 1976; Birnie & Paterson, 1991) and reactive Mn oxides, especially in suboxic aqueous systems such as the upper soil. The vernadite structure consists in the turbostratic stacking of layers of (Mn⁴⁺O₆)⁸⁻ octahedra connected through their edges and separated from each other by hydrated interlayer cations (e.g., Giovanoli, 1980; Chukhrov *et al.*, 1985; Jurgensen *et al.*, 2004; Villalobos *et al.*, 2006; Grangeon *et al.*, 2008). In natural settings, vernadite mainly has hexagonal layer symmetry (e.g., Marcus *et al.*, 2004; Manceau *et al.*, 2007, 2014; Bargar *et al.*, 2009). These layers may contain vacancies and/or isomorphic substitutions of Mn⁴⁺ by cations of lower

valence (*e.g.*, Mn^{3+} , Ni^{2+} , Co^{3+} ; Manceau *et al.*, 1997, 2014; Peacock & Sherman, 2007; Lanson *et al.*, 2008; Peña *et al.*, 2010; Grangeon *et al.*, 2014) which, together with the minute size of the layers (5–10 nm, *e.g.*, Hochella *et al.*, 2005; Bargar *et al.*, 2009), are responsible for the exceptional reactivity of vernadite. The possible contact with redox-sensitive species such as Mn^{2+} can induce structural changes whose nature and extent depend on solution pH and Mn^{2+} concentration. The first step of reaction between aqueous Mn^{2+} and layer Mn^{4+} , hereafter referred to as $\text{Mn}^{2+}_{(\text{aq})}$ and $\text{Mn}^{4+}_{(\text{s})}$, respectively, is (Elzinga, 2016):



where $\text{Mn}^{3+}_{(\text{s})}$ is layer Mn^{3+} . The $2\text{Mn}^{3+}_{(\text{s})}$ formed may further disproportionate, returning the system to its initial state. Alternatively, they may accumulate in the structure and induce a change in layer symmetry, from hexagonal to orthogonal, as observed experimentally at pH 9 when the molar ratio of Mn^{2+} to $\delta\text{-MnO}_2$ is lower than 0.05 (Zhao *et al.*, 2016). At lower pH and/or at higher Mn^{2+} concentration, a transformation to synthetic feitknechtite ($\beta\text{-MnOOH}$) is observed, following the reaction:



Feitknechtite, whose structure is built of layers of $(\text{Mn}^{3+}\text{O}_6)^{9-}$ octahedra, may upon ageing convert to manganite ($\gamma\text{-MnOOH}$) at pH 7.0–7.5 or to hausmannite (Mn_3O_4) at pH > 8 (Elzinga, 2011; Lefkowitz *et al.*, 2013; Lefkowitz & Elzinga, 2015).

The interplay between these structural transformations certainly influences the geochemical cycle of trace elements by modifying the reactivity of Mn oxides through a change in the nature and density of local crystal charges, but also by modifying the fate of trace elements initially adsorbed at the vernadite surface which, by analogy with the phyllo-manganate-to-tectomanganate transformation, could either be incorporated in the structure during transformation or released to solution (Atkins *et al.*, 2014, 2016; Grangeon *et al.*, 2015).

The present study aimed at contributing to a better understanding of the vernadite-to-feitknechtite transformation, and in particular at elucidating the reaction mechanisms at the crystal scale, using wet chemistry, X-ray diffraction (XRD) and transmission electron microscopy (TEM).

2. Materials and methods

2.1. Sample synthesis, sorption experiments and chemical analyses

The $\delta\text{-MnO}_2$ phase was synthesized using a redox method described elsewhere (Villalobos *et al.*, 2003). The average oxidation state (AOS) of Mn was determined by using a potentiometric method (Grangeon *et al.*, 2012). After synthesis, samples were Na-saturated through ten cycles of centrifugation, supernatant removal and resuspension in a

1 M NaCl solution. A last step of three similar cycles with de-ionized water instead of 1 M NaCl was then performed. The obtained $\delta\text{-MnO}_2$ paste was immediately freeze-dried, and the resulting powder was introduced in a N_2 -filled glove box where it was left for ~ 7 d. Eight aliquots of $\sim 20\text{--}40$ mg powder each were then introduced in previously weighted dialysis membranes. Filled membranes were weighted, filled with a solution containing 0.1 M NaCl and MnCl_2 at a concentration ranging 0–0.04 M, sealed, and introduced in 50 mL polypropylene tubes, to which 40.5–45 mL of the solution introduced in the dialysis membrane were added. Initial solution pH was 6.5. Tubes were sealed and left to agitate for 24 h. The dialysis membrane was then separated from the solution whose final pH ranged between 6.5 and 5.8. The solution was acidified to pH ~ 3 with HNO_3 and analyzed for its Mn^{2+} content using an ICP-AES (Horiba-Jobin Yvon Ultima 2). The magnitude of Mn^{2+} retention by $\delta\text{-MnO}_2$ was quantified using the R_D factor (Tournassat *et al.*, 2013) defined as:

$$R_D = \frac{C_{\text{init}} - C_{\text{aqu}}}{C_{\text{aqu}} \times R_{\text{SL}}},$$

where C_{init} is the initial concentration of Mn^{2+} , C_{aqu} the Mn^{2+} concentration in solution after equilibration, and R_{SL} is the solid-to-liquid ratio in the experiment (in kg L^{-1}).

The dialysis membrane was opened and the solid separated by filtration (0.01 μm cut-off) before being washed with bi-distilled water. An aliquot was freeze-dried for TEM analysis, whereas the remaining was sealed in a polyimide capillary for XRD analysis in the wet state within 48 h. Samples were labelled MndBi-XX, where XX is the Mn^{2+} loading (in mmol Mn^{2+} per g of dry $\delta\text{-MnO}_2$).

2.2. Synchrotron X-ray diffraction

The XRD experiments were carried out at station CRISTAL (SOLEIL synchrotron in Orsay, France), using an energy of 28.44 keV ($\lambda = 0.436 \text{ \AA}$) and an XPad hybrid pixel detector. Data were recorded over the 1.2–124.5° 2θ range with a total collection time of 30 min and processed with a specific software (Ounsi *et al.*, 2013). To ease comparison with previous studies (*e.g.*, Grangeon *et al.*, 2015, 2017), diffraction data will be expressed relative to the scattering vector q , where $q = [4 \times \pi \times \sin(\theta)]/\lambda$.

2.3. Transmission electron microscopy

The TEM experiments were carried out using a Philips CM20 microscope operated at 200 kV. Prior to observation, samples were embedded in epoxy resin, left to polymerize for 48 h in the dark and cut with an ultramicrotome (Reichert-Jung Ultra-cut E) equipped with a diamond knife. The ~ 80 nm thick sections were picked up on lacey carbon films loaded on Cu grids. In addition, to assess possible preparation-induced artefacts (use of an ultramicrotome), a sample identical to MndBi-0.5 (Table 1) was prepared according to the above described protocol, filtered and

Table 1. Experimental conditions of δ -MnO₂ contacts with Mn²⁺_(aq) and chemical characterization of reacted solids.

Sample	Mn AOS	Mn ²⁺ _{sorbed} (mmol g ⁻¹)	C _{aqu} (mmol L ⁻¹)	R _{SL} (g L ⁻¹)	log(R _D) (R _D in L kg ⁻¹)
MndBi-0	3.94 ± 0.05	n.d.	n.d.	0.65	n.d.
MndBi-0.5	3.80 ± 0.05	0.49	0.03	0.70	4.21
MndBi-1.5	n.d.	1.49	0.41	0.68	3.56
MndBi-3.3	3.54 ± 0.10	3.27	0.94	0.73	3.54
MndBi-8.6	n.d.	8.61	2.41	0.49	3.55
MndBi-11.3	n.d.	11.32	6.35	0.91	3.25
MndBi-28.1	3.62 ^a	28.05	12.15	0.64	3.36
MndBi-31.9	n.d.	31.90	20.47	0.55	3.19

Notes: Mn²⁺ loading at the end of the experiment (Mn²⁺_{sorbed}) is the number of mmol of Mn²⁺ sorbed per g of δ -MnO₂. n.d. stands for “not determined”.

^a May be subject to high uncertainties as a single measurement was performed.

re-suspended in alcohol. A drop of the obtained suspension was deposited on a Cu grid. Fast-Fourier transform (FFT) analysis of the micrographs was done with ImageJ (Schneider *et al.*, 2012).

3. Results and discussion

3.1. Determination of Mn²⁺ retention coefficient

An increase in Mn²⁺_(aq) resulted in a decrease in the Mn AOS of the final solids, suggesting retention of Mn²⁺ on δ -MnO₂ (Table 1). The R_D values (Table 1) compared well with previously reported values obtained after 8 d of equilibration (Fig. 1). The evolution of R_D as a function of Mn²⁺ concentration and pH (Murray, 1975) has, in some cases, been interpreted in terms of adsorption and quantified in surface-complexation models (Appelo & Postma, 1999; Tonkin *et al.*, 2004). Data interpreted with these models had a Mn²⁺ loading ranging from 1 to 12 Mn²⁺ per nm², depending on the dataset and on the specific surface area considered (value measured by the authors or 600 m² g⁻¹ as recommended by Tonkin *et al.*, 2004). Such loading roughly corresponds to MndBi-1.5 (1.5 Mn²⁺ per nm² assuming a specific surface area of 600 m² g⁻¹), which means that this sample and MndBi-0.5 could have been interpreted in the frame of these models. For other samples, the density of Mn²⁺ at the surface of δ -MnO₂ would be unrealistically high (up to 32 Mn²⁺ per nm² for MndBi-31.9), and interaction of Mn²⁺ with δ -MnO₂ most likely resulted in a structure transformation, which was studied by XRD.

3.2. Mineralogical evolution

Two sets of reflections could be observed in the XRD pattern of MndBi-0 (Fig. 2), which was not in contact with Mn²⁺, both being consistent with a nanocrystalline turbostratic phyllosilicate structure. The first set corresponded to broad and asymmetric reflections at 2.58 Å⁻¹, 4.42 Å⁻¹ and 5.11 Å⁻¹ (*d*-spacing of 2.44 Å, 1.42 Å and 1.23 Å). The reflection at 4.42 Å⁻¹ was shown to be indicative of layer symmetry: approximately symmetric when layer symmetry is hexagonal, but split in two distinct maxima when layer symmetry is orthogonal

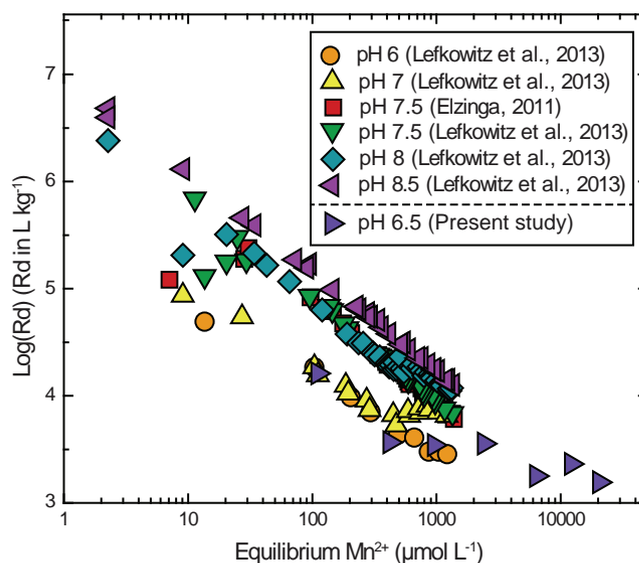


Fig. 1. Evolution of the Mn²⁺ retention coefficient as a function of Mn²⁺ concentration. Patterns are indicative of the experiment pH: circles and triangles pointing up, experiments conducted at pH 6 and pH 7, respectively; squares and triangles pointing down, pH 7.5; diamonds, pH 8; triangles pointing left, pH 8.5 (Elzinga, 2011; Lefkowitz *et al.*, 2013). Published data were collected with a contact time of 8 d. Present data (pH 6.5; 1 d contact time) are shown as triangles pointing right.

(Drits *et al.*, 2007). Here, it was symmetric despite the presence of a shoulder on the low-*q* side, related to a basal reflection (see below), and layer symmetry was hexagonal. For consistency with previous studies (*e.g.*, Lanson *et al.*, 2000, 2008; Webb *et al.*, 2005; Drits *et al.*, 2007; Grangeon *et al.*, 2010, 2012; Zhu *et al.*, 2010; Manceau *et al.*, 2013), bands were indexed with an orthogonal symmetry system ($a = \sqrt{3} \times b$, $\gamma = 90^\circ$) and the three maxima were indexed as (11, 20), (31, 02) and (22, 40) *hk* band, respectively. Reflections of the second set were sharp and symmetric and formed a rational series of 00*l* reflections at 0.62 Å⁻¹, 1.24 Å⁻¹ and 1.85 Å⁻¹ (10.15 Å, 5.08 Å and 3.39 Å). The layer-to-layer distance, inferred from the position of the 003 at 1.85 Å⁻¹, was 10.17 Å. When increasing Mn²⁺ loading up to 1.5 mmol g⁻¹ (MndBi-1.5), the intensity of the 00*l* reflections decreased, but the position remained essentially unchanged and

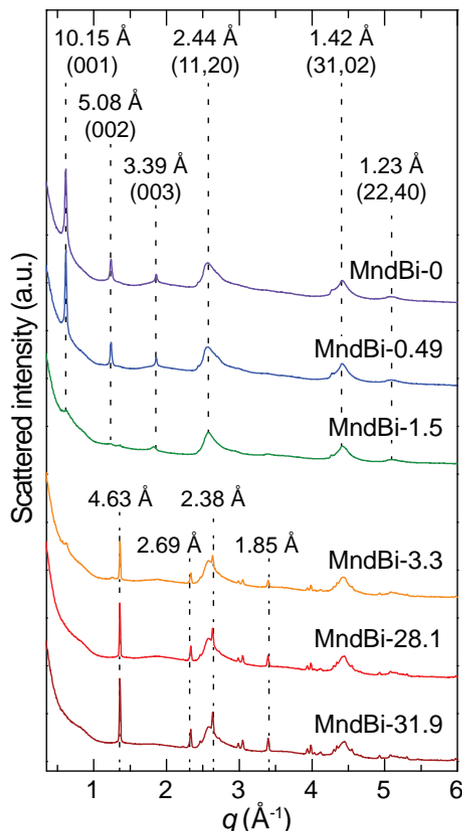


Fig. 2. From top to bottom, XRD patterns of MndBi-0, MndBi-0.5, MndBi-1.5, MndBi-3.3, MndBi 28.1 and MndBi-31.9. Dashed lines and dashed-dotted lines point out to the presence of diffraction maxima of δ -MnO₂ and synthetic feitknechtite, respectively.

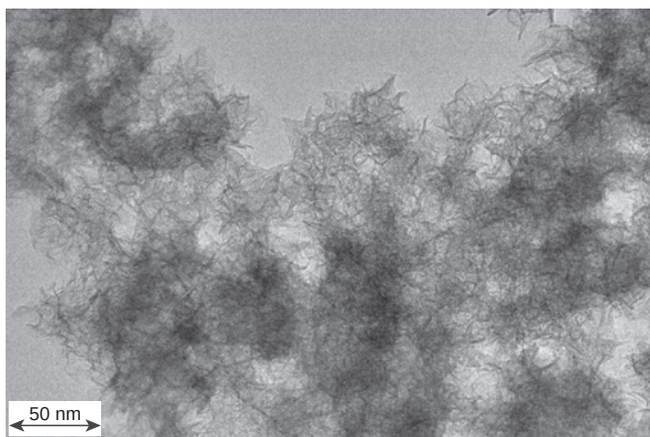


Fig. 3. Transmission electron microscope (TEM) observation of type 1 crystals from MndBi-0.

hk bands were unaffected. The shoulder on the low-*q* side of the (31, 02) band weakened, supporting its indexation as the 006 reflection. Note the presence of a weak peak at $\sim 1.36 \text{ \AA}^{-1}$ ($\sim 4.6 \text{ \AA}$) in the XRD pattern of MndBi-1.5, which could be attributed to synthetic feitknechtite (see below). Weakening of 00*l* reflections can be interpreted as a loss of parallelism between adjacent layers and was attributed to the incorporation of Mn³⁺_(s) according to the

comproportionation mechanism described above. Indeed, Mn³⁺ octahedra are distorted by the Jahn–Teller effect and their presence within the octahedral layer induces strains that ultimately lead to layer kinking (Webb *et al.*, 2005; Atkins *et al.*, 2014; Grangeon *et al.*, 2014) and subsequent loss of parallelism between layers. In addition to *hk* bands of δ -MnO₂, the XRD pattern of MndBi-3.3 (3.3 Mn²⁺ per nm²) had additional reflections, *e.g.*, at 1.36 \AA^{-1} , 2.34 \AA^{-1} , 2.64 \AA^{-1} and 3.40 \AA^{-1} (4.63 Å, 2.69 Å, 2.38 Å, and 1.85 Å), assigned to synthetic feitknechtite (Bricker, 1965). As the latter reflections became more intense with increasing Mn²⁺ loading, up to $31.90 \text{ mmol g}^{-1}$ (MndBi-31.9), the proportion of synthetic feitknechtite increased with Mn²⁺ loading.

Such an evolution was consistent with previous studies (Elzinga, 2011, 2016; Lefkowitz *et al.*, 2013; Lefkowitz & Elzinga, 2015). In the following section, the mechanisms of transformation were studied at the crystal scale, hypothesizing that an increased concentration of Mn²⁺ led to an increased conversion of δ -MnO₂ to synthetic feitknechtite without significant dissolution of δ -MnO₂. In other words, it was hypothesized that the structural and morphological evolution of samples following their contact with a given concentration of Mn²⁺_(aq) underwent intermediate recrystallization steps similar to those observed during *post-mortem* analysis of samples in contact with lower concentrations of Mn²⁺_(aq). This hypothesis is supported by previous TEM observations, from which it was concluded that there was a lack of significant dissolution/precipitation during the recrystallization of δ -MnO₂ when in contact with aqueous MnSO₄ (Tu *et al.*, 1994). Absence of δ -MnO₂ dissolution is further substantiated by the study of Elzinga & Kustka (2015) showing that the first steps of transformation involves solely comproportionation and disproportionation reactions induced by interfacial electron transfer between adsorbed Mn²⁺ and Mn⁴⁺_(s). Experimental validation of this hypothesis would require performing a kinetic experiment at high Mn²⁺_(aq) concentration to check whether recrystallization steps observed as a function of Mn²⁺ concentration are similar to those occurring as a function of time. Such a procedure was applied to the study of the δ -MnO₂ to synthetic todorokite transformation (Atkins *et al.*, 2014, 2016), but was not possible in the present case because of reaction kinetics. The *R_D* values obtained in the present study after 1 d of contact time were identical to those reported previously for a contact time of 8 d (Fig. 1). Steady-state was thus reached in less than 1 d of contact time, consistent with previous observations showing that feitknechtite is formed in less than 20 min (Johnson *et al.*, 2016) and that sorption of Mn²⁺ at the δ -MnO₂ surface is completed within less than 1 s (Fendorf *et al.*, 1993).

3.3. Morphological evolution during δ -MnO₂ to synthetic feitknechtite transformation

Sample MndBi-0 consisted of crystals $\sim 10 \text{ nm}$ in their largest dimension (presumably the layer plane) and $\sim 1 \text{ nm}$ perpendicular (Fig. 3), henceforth denominated

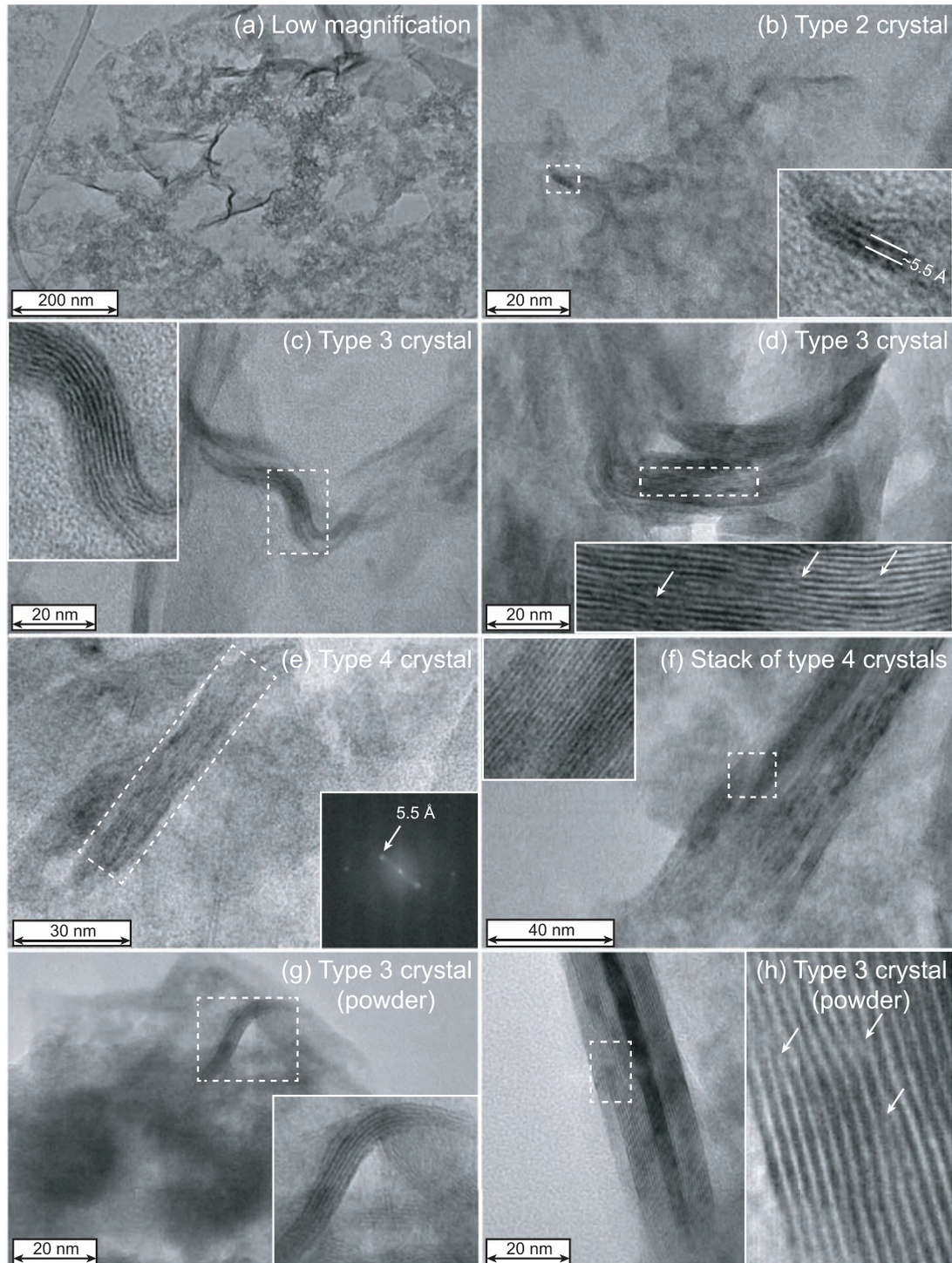


Fig. 4. TEM observation of $\text{Mn}_\delta\text{Bi-0.5}$. (a) Low-magnification view; (b) type-2 crystal; (c) and (d) type-3 crystals; (e) type-4 crystal; (f) crystal built from the stacking of type-4 crystals; (g) and (h) type-3 crystals deposited on the Cu grid from a drop of suspension instead of microtome preparation. Insets in (b), (c), (d), (f), (g) and (h) are enlarged views of regions delimited with dotted squares. In (d) and (h), arrows point to the presence of dislocations. The inset in (e) is a fast-Fourier transform (FFT) analysis of the region shown with a dotted square.

type-1 crystals. The crystal size in the layer plane was in good qualitative agreement with the XRD data, in which broad hk bands could be observed, indicative of crystallites $\sim 5\text{--}10$ nm in the layer plane (Lanson *et al.*, 2008; Grangeon *et al.*, 2012; Manceau *et al.*, 2013). On

the other hand, the small crystal size perpendicular to the layer plane contrasted with the sharp $00l$ reflections. This apparent contradiction could be concealed assuming that TEM sample preparation procedure has delaminated $\delta\text{-MnO}_2$ crystals.

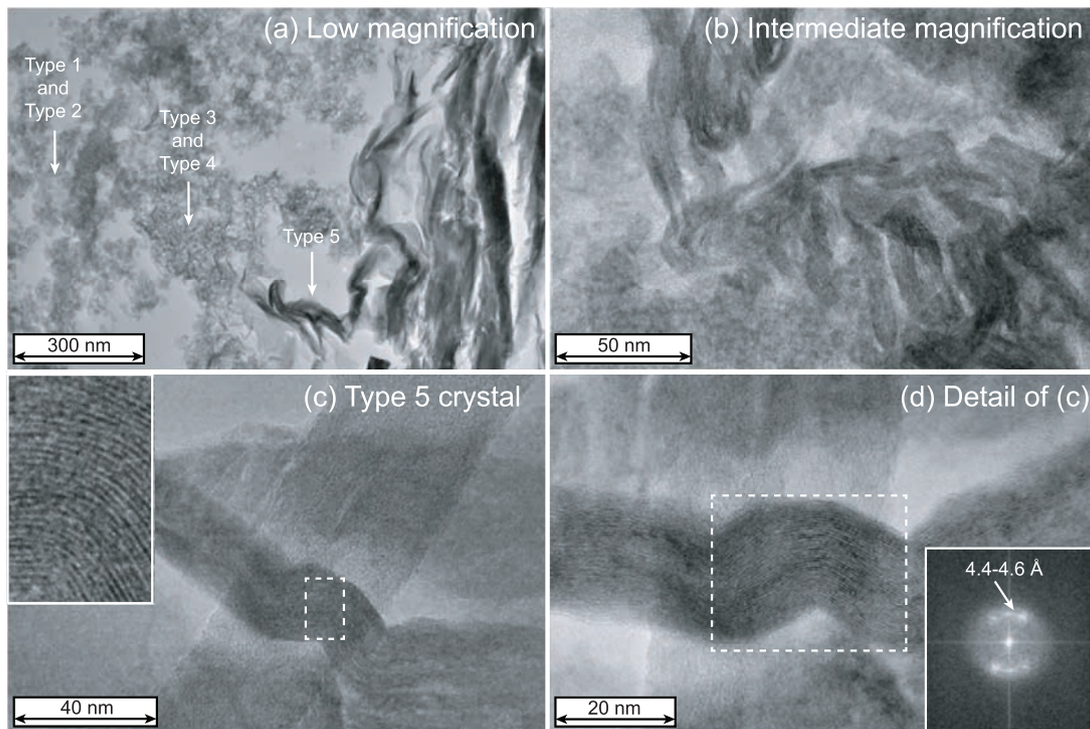


Fig. 5. TEM observation of MndBi-1.5. (a) Low-magnification view with the different crystal morphologies indicated with arrows; (b) type-5 crystals at intermediate magnification; (c) type-5 crystal exhibiting layer bending. The area shown with a dotted square in (c) is enlarged in the inset. (d) A detailed view of (c) and the inset is a FFT analysis of the region shown with a dotted square.

Observation of MndBi-0.5 evinced that a Mn^{2+} loading as low as 0.49 mmol g^{-1} led to the formation of crystals with contrasting morphologies and sizes, ranging ~ 10 – 200 nm in the layer plane (Fig. 4a). In addition to type-1 crystals, three types of such crystals were identified:

- Type-2 crystals had a size of $\sim 10 \text{ nm}$ in the layer plane, ~ 2 – 3 nm perpendicular, and were bent (Fig. 4b), probably as a result of the incorporation of $\text{Mn}^{3+}_{(s)}$ in layers containing mainly Mn^{4+} . The layer-to-layer distance was $\sim 5.5 \text{ \AA}$, typical for dehydrated phyllosulfates (Cygan *et al.*, 2012; Wegorzewski *et al.*, 2015). Dehydration was possibly induced by sample freeze-drying or by the secondary vacuum in the TEM. For type-1 and type-2 crystals having similar lateral dimensions, it is proposed that the latter formed from the coherent stacking of the former;
- Type-3 crystals were $\sim 50 \text{ nm}$ in the layer plane, and were built of ~ 10 – 20 layers stacked parallel to each other, with a layer-to-layer distance of $\sim 5.5 \text{ \AA}$ (Fig. 4c, d). As type-2 crystals, they were bent. Images of the lateral terminations of type-3 crystals showed that they consisted of stacked type-2 crystals (Fig. 4c). Layer dislocations were observed also on some crystals (Fig. 4d), suggesting that type-2 crystals connected also within the layer plane to form type-3 crystals. Most likely, the type-3 crystals thus originated from the oriented aggregation of type-2 crystals, both within the layer plane and perpendicular to it;
- Type 4 crystals had lath-like shape, dimensions of ~ 50 – 100 nm in the layer plane and ~ 10 – 20 nm perpendicular (Fig. 4e), layer-to-layer distance of

5.5 \AA (inset in Fig. 4e), and were rarely found as stacks (Fig. 4f). It is speculated that they resulted from the same aggregation mechanisms proposed for the type-3 crystals, but with fewer defects between connected type-2 crystals and/or a specific distribution of $\text{Mn}^{3+}_{(s)}$ so as to minimize layer strains and bending. Note that layer bending and dislocations observed in type-3 crystals cannot be related to sample preparation, as crystals deposited from a drop of a $\delta\text{-MnO}_2$ suspension exhibited similar microstructural features (Fig. 4g, h).

These observations demonstrate that the surface of $\delta\text{-MnO}_2$ is modified at loadings as low as 0.5 Mn^{2+} per nm^2 . As a consequence, surface complexation models, which rely on a reversible sorption hypothesis (Tournassat *et al.*, 2013), may not be suited to describe Mn^{2+} interaction with $\delta\text{-MnO}_2$.

All types of crystals found in samples with lower Mn^{2+} loadings were observed in MndBi-1.5 that contained also larger crystals (type-5), ~ 100 – 150 nm in the layer plane and ~ 20 – 30 nm perpendicular (Fig. 5a). In contrast, XRD indicated that MndBi-1.5 consisted mainly of isolated $\delta\text{-MnO}_2$ nanosheets. However, type-5 crystals were bent (Fig. 5b). Bending did not involve layer disruption (Fig. 5c), but likely led to variable layer-to-layer distance, best quantified from FFT analysis of the images (Fig. 5d). This analysis showed that layer-to-layer distance varied within a given crystal, being equal to 4.4 – 4.6 \AA . Both layer bending and layer-to-layer distance heterogeneity within a given crystal reduced crystallite size within the layer plane and led to vanishing of $00l$ reflection.

Finally, MndBi-28.1 contained, in addition to all previously mentioned types of crystals, type-6 crystals (Fig. 6) whose selected-area electron diffraction (SAED) patterns were typical for synthetic feitknechtite (Fig. 6c). These crystals were up to $\sim 100 \text{ nm} \times 500 \text{ nm}$ and were built of the stacking of laths resembling type-5 crystals but free of stacking defects and dislocations. These laths were not simply aligned, but were connected (Fig. 6d, e), as shown by both the observation of interference fringes (inset in Fig. 6e) and the FFT image analysis showing a homogeneous layer-to-layer distance (4.4 \AA , inset in Fig. 6e). It is proposed that type-6 crystals formed from the oriented aggregation, perpendicular to the layer plane, of type-5 crystals. The presence of dislocations in type-5 crystals is thought to have inhibited their oriented aggregation. Bending was not observed either in type-6 crystals, possibly because all $\text{Mn}^{4+}_{(s)}$ was converted to $\text{Mn}^{3+}_{(s)}$, allowing for their specific distribution to accommodate their Jahn–Teller distortion. No Mn^{2+} -rich phase could be detected with SAED, despite a molar ratio of sorbed Mn^{2+} to structural $\delta\text{-MnO}_2$ Mn of 3 (assuming a defect-free $\delta\text{-MnO}_2$ and a molar mass of 119 g mol^{-1} , Giovanoli *et al.*, 1970). Consequently, it was concluded that part of the Mn^{2+} was sorbed by $\delta\text{-MnO}_2$ and/or synthetic feitknechtite, or precipitated as an amorphous phase.

3.4. The role of oriented aggregation in nanoparticle growth

The present description of the vernadite-to-feitknechtite transformation, from macroscopic to crystal scales, is consistent with our current understanding of this transformation (Elzinga, 2011, 2016; Lefkowitz *et al.*, 2013; Lefkowitz & Elzinga, 2015). One of the key findings is the major role of oriented aggregation in the nucleation and growth of feitknechtite from $\delta\text{-MnO}_2$ precursors. This process is increasingly recognized as one of the main processes responsible for nanoparticle growth (Penn, 2004; Niederberger & Colfen, 2006; Yuwono *et al.*, 2010): carbonates (Shen *et al.*, 2006), iron (Jia & Gao, 2008a and b) and manganese (Portehault *et al.*, 2007) oxides. It was also shown to be responsible for the phase transformations of nanoparticle precursors (*e.g.*, Hockridge *et al.*, 2009). Recently, oriented aggregation was shown to be involved in the formation of synthetic todorokite and cryptomelane from a $\delta\text{-MnO}_2$ precursor (Chen *et al.*, 1986; Atkins *et al.*, 2014, 2016; Grangeon *et al.*, 2015) and in the change of $\delta\text{-MnO}_2$ layer symmetry, from hexagonal to orthogonal (Zhao *et al.*, 2016). It is thus possible to speculate that Mn oxides found in surficial environments were to a large extent formed from vernadite precursors (Bodeř *et al.*, 2007; Xu *et al.*, 2010), possibly through oriented aggregation mechanisms.

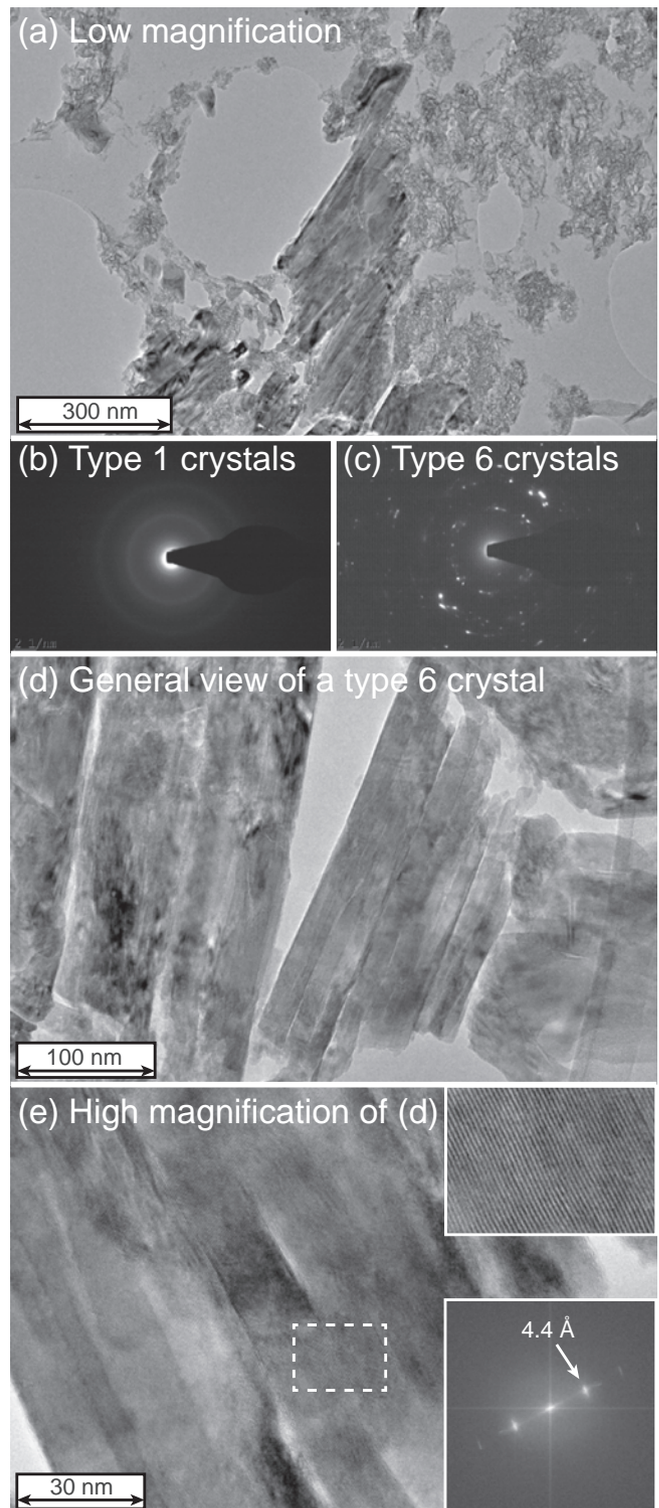


Fig. 6. TEM observation of MndBi-28.1. (a) Low-magnification view; (b) and (c) SAED patterns collected on the smallest crystals (top right in (a)) and the biggest crystals (centre in (a)), respectively; (d) and (e) type-6 crystal. The FFT analysis shown as an inset at the bottom right of (e) was calculated over the whole image while the inset at the top right is an enlarged view of the area delimited with a dotted square.

4. Concluding remarks: environmental implications

Vernadite with hexagonal layer symmetry is a sink for many trace elements in the environment. In natural settings $\text{Mn}^{2+}_{(\text{aq})}$ is abundant, such as in lake and marine water columns and in sediments (Burdige, 1993), interactions between vernadite and $\text{Mn}^{2+}_{(\text{aq})}$ may induce morphological and mineralogical changes that could impact its reactivity and the fate of trace elements sorbed at its surface. Two main structural transformations were identified during reaction of $\delta\text{-MnO}_2$ (synthetic vernadite) with $\text{Mn}^{2+}_{(\text{aq})}$. At a low Mn^{2+} to $\delta\text{-MnO}_2$ ratio ($<3.27 \text{ mmol g}^{-1}$), $\delta\text{-MnO}_2$ crystal size increased with $\text{Mn}^{2+}_{(\text{aq})}$. At a Mn^{2+} to $\delta\text{-MnO}_2$ ratio of 3.27 mmol g^{-1} or higher, $\delta\text{-MnO}_2$ was converted to synthetic feitknechtite, and the size of synthetic feitknechtite crystals was observed to increase with $\text{Mn}^{2+}_{(\text{aq})}$. $\delta\text{-MnO}_2$ with orthogonal layer symmetry was not detected as a transformation intermediate, possibly because experiments were conducted at slightly acidic pH (Zhao *et al.*, 2016).

When crystal size increases, the specific surface area, and especially the contribution of particle edges to this surface, decreases. Vernadite edge sites being reactive (Simanova *et al.*, 2015), the increase in vernadite crystal size when in contact with $\text{Mn}^{2+}_{(\text{aq})}$, even at low concentrations, can be detrimental to its trace-metal scavenging ability. By analogy, the same is expected for feitknechtite at high Mn^{2+} to $\delta\text{-MnO}_2$ ratio. The impact of the vernadite-to-feitknechtite transformation on the fate of trace elements remains unclear, however. The lack of any significant dissolution of $\delta\text{-MnO}_2$ potentially minimizes the release of trace metal elements along the reaction pathway, but their possible incorporation in the newly formed feitknechtite structure remains undocumented. Unlike tunnel structures, feitknechtite has been observed seldom in natural settings, probably because of its limited chemical stability and high sensitivity to fluctuation in redox conditions, which lead to its back-conversion to vernadite (Bargar *et al.*, 2005). Consequently, no data are currently available to evaluate the affinity of trace elements for feitknechtite in natural settings. Similar to tunnel structures (Liu *et al.*, 2004; Kumagai *et al.*, 2005; Cui *et al.*, 2011; Atkins *et al.*, 2014), additional insights into vernadite reactivity in suboxic systems and its impact on the fate of trace metals could probably be sought experimentally as feitknechtite is frequently observed in laboratory experiments mimicking natural conditions (Hem *et al.*, 1982; Murray *et al.*, 1985; Wang *et al.*, 2015).

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