The role of biogenic Fe-Mn oxides formed *in situ* for arsenic oxidation and adsorption in aquatic ecosystems

Yaohui Bai, Tingting Yang, Jinsong Liang, Jiuhui Qu

A Key Laboratory of Drinking Water Science and Technology, Research Center for Eco-Environmental Sciences, Chinese Academy of Sciences, Beijing 100085, China

B University of Chinese Academy of Sciences, Beijing 100049, China

**Abstract**

As(III & V), Mn(II), and Fe(II) may occur simultaneously in some groundwater and surface water. Studying their redox reactions and interactions is essential to unravel the biogeochemical cycles of these metal ions in aquatic ecosystems and to find effective methods to remove them simultaneously in drinking water treatment. Here, the formation of biogenic Fe-Mn oxides (BFMO, defined as a mixture of biogenic Mn oxide (BMO) and Fe oxide) as well as its oxidation and adsorption of As in a Fe(II)-Mn(II)-As(III & V)-Mn-oxidizing microbe (Pseudomonas sp. QJX-1) system was investigated. Batch experiments and structure characterization revealed that the BFMO was formed via a sequential precipitation of Fe oxide and BMO. The first formed Fe oxide was identified as FeOOH (lepidocrocite) and the latter formed BMO was identified as MnO₂ (similar to hexagonal birnessite). In the BFMO mixture, the BMO part was mainly responsible for As(III) oxidation, and the Fe oxide part dominated As adsorption. Remarkably, the BMO could oxidize Fe(II) to form FeOOH, which may improve As adsorption. The optimum Mn(II)/Fe(II) ratio for As removal was approximately 1:3 (mol/mol). Taken together, in Fe(II)-Mn(II)-As(III & V)-Mn-oxidizing microbe ecosystems, the *in situ* formation of BFMO could eliminate or decrease Fe(II), Mn(II), and As(III & V) species simultaneously. Therefore, based on this study, new approaches may be developed for As removal from water containing high concentrations of Fe(II) and Mn(II).

1. Introduction

In groundwater and in some surface water, soluble Fe(II) and Mn(II) often occur together (Huang et al., 2012; Postawa and Hayes, 2013), and Fe and Mn oxides are produced by oxidative precipitation under aerated conditions. Fe oxide is mainly formed by Fe(II) chemical oxidation (Kappler and Straub, 2005), and Mn oxide is formed via biological Mn(II) oxidation. Accordingly, Fe-Mn oxides are often formed *in situ* in natural aquatic ecosystems as well as in groundwater treatment systems (Bai et al., 2013). Extensive studies have suggested that biogenic Mn oxide (BMO) is an efficient sorbent and oxidant due to its highly charged surface and oxidizing power (Hansel et al., 2011; Tebo et al., 2010); and synthetic Fe oxide is a powerful sorbent that is currently used in drinking water treatment processes (Faust, 1998). However, the oxidation and adsorption characteristics of the biogenic Fe-Mn oxides (defined as BMO combined with Fe oxide, BFMO) formed *in situ* remain largely unknown.

Arsenic (As) is a ubiquitous contaminant that jeopardizes water quality, and its trivalent species has higher toxic effects (Gebel, 1997). High concentrations of As are found in some surface water and groundwater along with high Fe(II) and Mn(II) concentrations, and all of these contaminants present a threat to public health and aquatic ecology (Davolos and Pietrangeli, 2013). For treatment, the desirable solution is to eliminate or decrease these three ions simultaneously from contaminated water. Based on this, some treatment methods/processes was developed, for example, aeration together with chlorination or ozonation followed by filtration can be used to remove Fe(II) and Mn(II), which would also remove As species (depending on the As/Fe ratio) (Nishimura and Umetsu, 2001). Greensand filters (Viraraghavan et al., 1999) and household-scale sand filters (Berg et al., 2006; Voegelin et al., 2014) have also been shown to remove them simultaneously. Here we provide another treatment concept, by means of strengthening the formation of BFMO in the treatment system. The following hypothesis based on the oxidation and adsorption properties of BMO (Murray...
et al., 2005; Wu et al., 2005) and Fe oxide (Fig. 1) was proposed: When a Mn-oxidizing microbe such as *Pseudomonas* sp. QXJ-1 is introduced, soluble Mn(II) will be oxidized to insoluble Mn(IV) oxide (BMO), which will mix with insoluble Fe oxide produced from chemical Fe(II) oxidation in an aerated environment to form BFMO. The BFMO formed in situ will then oxidize and adsorb As species. It was assumed that BMO oxidize (and adsorb?) As and that Fe oxide adsorb As. Meanwhile, As oxidation may lead to the reduction of Mn oxide to Mn(II). However, Mn(II) will be reoxidized biologically to BMO. With this cycle, the Fe(II), Mn(II), and As(III&V) species may be efficiently removed from the water.

In this study, the hypothesis was experimentally verified, and the valence shifts of Fe, Mn, and As during the reaction, the respective roles of Mn oxide and Fe oxide in As oxidation and adsorption, and the interaction between Fe oxide and Mn oxide were also clarified. Therefore, the findings from this study may not only shed light on the biogeochemical cycles of three elements in natural or engineered ecosystems but also may provide a new method to treat water containing Mn(II), Fe(II), and As(III&V) species.

2. Materials and methods

2.1. Mn(II)-oxidizing bacterial strain

A Mn(II)-oxidizing bacterium was isolated from the soil of a Mn mine, located in Xiangtan City, Hunan Province, China. According to the physicochemical and 16S rRNA characterizations, the bacterium was identified and named as *Pseudomonas* sp. QXJ-1 (GeneBank accession number, KM242057). QXJ-1 is a Gram-negative, rod strain with dimensions of 1–2.1 μm in length and 0.5–0.9 μm in width. A culture of the strain has been deposited in the China General Microorganism Culture Center (CGMCC; accession number, 6630).

2.2. Enrichment of inoculated bacteria

Peptone–yeast extract–glucose (PYG) medium, described by Adams and Ghiorse (1985), was modified slightly and used in the experiments. Each liter of PYG contained 0.25 g of peptone, 0.25 g of yeast extract, 0.25 g of glucose, 0.5 g of MgSO4·7H2O, and 0.06 g of CaCl2·2H2O. The pH of the PYG medium was adjusted to 7.5 with 1 M NaOH before autoclaving. Approximately 15 mM N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid (HEPES) was used as the buffer in the PYG medium.

*Pseudomonas* sp. QXJ-1 was inoculated in several 250-ml flasks containing 100 ml of PYG media and incubated at 30 °C and 170 rpm in a rotary shaker until the bacteria reached the logarithmic phase of growth (approximately 48 h). The bacterial suspensions were used immediately for the following oxidation and adsorption experiments.

2.3. BFMO formation and its oxidation and adsorption for As species

First, BFMO formation and its removal of As with different initial Mn/Fe ratios (mol/mol) were investigated to determine the optimum ratio. The experiment used a series of 500-ml Erlenmeyer flasks as batch reactors. Each flask contained 200 ml of PYG medium containing 100 μM Mn(II) (prepared with MnCl2), 10 μM As(III&V), prepared with sodium arsenite (NaAsO2) and sodium arsenate (NaAsO4·2H2O), respectively, and different initial concentrations of Fe(II) (prepared with FeSO4·7H2O). The initial inoculum sample had an optical density at 600 nm, corresponding to less than 0.1 units. Flasks without the bacterial inoculum were used as negative controls under the same conditions.

In this study, the hypothesis was experimentally verified, and the valence shifts of Fe, Mn, and As during the reaction, the respective roles of Mn oxide and Fe oxide in As oxidation and adsorption, and the interaction between Fe oxide and Mn oxide were also clarified. Therefore, the findings from this study may not only shed light on the biogeochemical cycles of three elements in natural or engineered ecosystems but also may provide a new method to treat water containing Mn(II), Fe(II), and As(III&V) species.
scanning electron microscopy (FE-SEM) coupled with energy-dispersive X-ray analysis (EDX) were used to investigate the morphology, size, and element distribution of the BMO, Fe oxide, and BFMO. The media producing BMO, Fe oxide, and BFMO (described above) were incubated for 72 h. Suspensions were collected, centrifuged (5000 x g) and washed twice with sterile Milli-Q water to remove the organic matters and salts in the media. Finally, the collected precipitates were resuspended in 1 ml of Milli-Q water. The samples were divided into two aliquots. One was mounted on double-sided carbon tape and sputter-coated with a Pt/Pd film prior to imaging with a FE-SEM/EDX analyzer (Hitachi SU-8020). The other was imaged with a Hitachi H-7700 transmission electron microscope. For X-ray Powder Diffraction (XRD) analysis, BMO, Fe oxide and BFMO precipitates were collected by centrifugation of incubated suspensions (1–2L, 72 h) and then dried with a freeze drier. The dried samples were ground into powder using pestle and mortar. Finally the pressed powder disk was prepared and then analyzed using an X’Pert PRO MPD X-ray spectrometer (Netherlands) with Cu-Kα(αlpha) irradiation, PIKcel detector, generator voltage of 40 kV and tube current of 40 mA. The data were collected at a 0.712°/min of scanning speed in the angle range from 10° to 80° for BMO, and from 5° to 90° for Fe oxide and BFMO.

To further understand the valence shifts of As, Fe, and Mn in the experiment as well as the coordination of As associated with the BMO and BFMO, the lyophilized powder pellets of BMO, Fe oxide, BFMO, and their As adsorption samples (72 h) were analyzed at the Beijing Synchrotron Radiation Facility (BSRF), China, for X-ray absorption near-edge structure (XANES) and K-edge extended X-ray absorption fine structure (EXAFS) analyses. Eight reference compounds including MnO, MnO₂, MnO₃, Mn₂O₃, Fe₂O₃, FeO₂, FeO₄, Na₂AsO₂, and Na₃AsO₄ 12H₂O (purchased from Sigma-Aldrich, USA or URCHEM, China) were simultaneously analyzed for use as model spectra to assist interpretation of spectra of BMO, Fe oxide, BFMO, and their As adsorption samples. Samples were measured in transmission or fluorescence mode (used for As analysis due to its low concentration) at room temperature on wiggler beamline 1W1B of BSRF. The Lytle detector was used. The beam spot was focused to 2 x 2 mm² by the toroidal mirror. A fixed-exit Si (1 1 1) double crystal monochromator was used and the harmonic content of the monochromer beam was rejected by detuning the double crystal monochromator. An energy range of -2000 to 1000 eV from the K-edge of Mn, Fe, and As was used for EXAFS data collection. Standard EXAFS data reduction procedures were carried out using the program package IFEFFIT (Newville, 2001). The K²-weighted χ(k) was Fourier-transformed into R space in the k ranges of 2.6–11.0 Å⁻¹ for Fe K, 2.8–11.0 Å⁻¹ for Mn K, and 2.4–12.4 Å⁻¹ for As K with a Hanning function window. Shell fitting was adapted for fitting the Fourier-transformed K²-weighted (R space) Mn, Fe, and As K-edge EXAFS spectra. The goodness of fit was evaluated by R-factor minimization.

3. Results

3.1. In situ formation of BMO by Pseudomonas sp. QJX-1

The crucial step of BFMO formation is the generation of BMO by Pseudomonas sp. QJX-1. So, the characteristics of Mn(II) oxidation by QJX-1 under the optimum temperature (30 °C) and pH (7.5) were investigated, as shown in Supplementary Fig. S1. The results showed that QJX-1 catalyzed the slow transformation of Mn(II) in the logarithmic growth phase. However, after the logarithmic growth phase and during the stagnant growth phase, Mn(II) was oxidized rapidly. Meanwhile, a brownish black precipitate (i.e., BMO) was produced.

3.2. In situ formation of BFMO and its capacity for As removal

Different concentrations of Fe(II) were added to the PYG media containing Mn(II) and As species, and QJX-1 was subsequently added. In the initial phase of the experiment, due to the rotation, the Fe(II) ions in the media had sufficient contact with oxygen. Therefore, it was observed that the dark yellow precipitation of Fe oxide occurred within minutes. Then, the BMO was produced by QJX-1 and mixed with Fe oxide to form the BFMO. Chemical analysis of Fe(II) and Mn(II) oxidation (Fig. S2) revealed that more than 95% of Fe(II) and Mn(II) were oxidized, except in the media containing As(III) or As(V) at a 1:9 Mn(II)/Fe(II) ratio. Fig. 2 shows that the total As removal increased along with the increase of Fe(II) concentration, implying that Fe oxide plays a dominant role in As removal. In general, As(V) removal by the BFMO was better than As(III) removal. The optimum Mn(II)/Fe(II) ratio for As removal was determined to be approximately 1:3 (mol/mol).

3.3. Oxidation and adsorption of As species by the BMO, Fe oxide, and BFMO formed in situ

The oxidation and adsorption characteristics of As species among the BFMO formed in situ (1:3 Mn(II)/Fe(II) ratio), BMO, and Fe oxide were compared in order to identify the respective roles of the BMO and Fe oxide parts in the BFMO. In all reaction systems, the buffers used in the culture media ensured little pH variation, from an initial pH of 7.50 to 7.33–7.54 (Table S1) at the end of the experiment. Therefore, the chemical precipitation of Mn and Fe oxides induced by alkaline changes were excluded, and Mn(II) oxidation and Fe(II) oxidation were attributed to the biological role of QJX-1 and the chemical role of oxygen, respectively. Fig. 3a–b shows the formation of BMO and its chemical role on As(III) and V). The results from the media containing only QJX-1 and As species (denoted as control in the figure) demonstrated that QJX-1 had negligible effects on the oxidation and adsorption of As species. Hence, As(III) oxidation (i.e., 97.5 ± 0.62% of As(III)) was oxidized to As(V) within 48 h in the media containing QJX-1, Mn(II), and As(III) was only attributed to the BMO formed in situ (Fig. 3a). However, only small portions of As(III) and As(V) were adsorbed on the surface of the formed BMO (<5%) (Fig. 3a–b). In contrast to the BMO, Fe oxide formed in situ could rapidly (within the minute scale) adsorb As(III) or As(V) on its surface (more than 92% within 1 h) (Fig. 3c–d). The BFMO formed in situ integrated the chemical characteristics of the BMO and Fe oxide; it could not only oxidize but also adsorb the As species (Fig. 3e–f). It was remarkable that during Mn(II) oxidation (later than the chemical oxidation of Fe(II)), the formed BMO could further oxidize the remaining Fe(II) in solution to Fe oxide. Moreover, the newly formed Fe oxide was able to adsorb As again, resulting in a decrease of the total As during the experiment.

3.4. Variation of ζ-potential and particle size

The variations of ζ-potential during the formation of BMO, Fe oxide, and BFMO as well as their adsorption of As species were measured (Fig. 4). The presence of lipopolysaccharides on the Gram-negative bacterial surface provides negatively charged areas for absorbing cationic species (Yang et al., 2015). Hence, the ζ-potential decreased rapidly with the fast growth of QJX-1. The presence of Mn(II) increased the ζ-potential of system, probably due to that bacteria adsorbed some Mn(II) on the surface. When As species were introduced, As(III) existed primarily as H₂AsO₃⁻, an uncharged species, and As(V) was present mainly as HAsO₄²⁻ and smaller amounts of H₂AsO₄ which led to a small decrease of the ζ-potential. When adding Fe(II) to the media, along with the formation of Fe
Samples were collected at 72 h. Data points represent the mean value (three replicates) gated using K-edge XANES and EXAFS analyses. Fig. 6a shows that the solid phases of the BFMO, BMO, and Fe oxide were investigated using K-edge XANES and EXAFS analyses. The initial concentrations of As(III) and As(V) were 10 μM, and the initial Mn(II) concentration was 5 mg/L. Different initial Fe(II) concentrations (5 mg/L, 15 mg/L, 25 mg/L, 35 mg/L, 45 mg/L) were used in the experiment to explore the optimum Mn(II)/Fe(II) ratios for total As removal. Samples were collected at 72 h. Data points represent the mean value (three replicates) ± standard deviation.

Due to the low concentration of As in the reaction systems, the zeta-potential decreased to the same level as media containing bacteria alone. Further, when the BMO was produced, the zeta-potential value had a significant increase due to the formation of MnOx on the bacterial surface.

During the formation of the BMO and Fe oxide separately as well as the BFMO, the suspended particle sizes of each of them all exhibited a gradient increase against the reaction time (before precipitation) (Fig. S3). The maximum suspended BFMO size (370–400 nm) was between those of the BMO (700–800 nm) and Fe oxide (330 nm).

### 3.5. TEM, FE-SEM/EDX, and XRD analysis of BMO, Fe oxide, and BFMO

TEM (Fig. S4) and SEM (Fig. S5) analyses of the formed BMO, Fe oxide, and BFMO precipitates (72 h) show they all had irregular morphology. In general, the sizes of precipitates (in μm scale) were BMO > BFMO > Fe oxide, which is consistent with the comparison of their suspended particle sizes (Fig. S3). EDX results (Fig. S5) indicate that the formed BFMO was not a homogenous compound as Fe/Mn ratios varied in different parts of BFMO. For XRD analysis, the XRD pattern of BMO showed (Fig. 5) that two reflections at 2.46 Å and 1.41 Å (d spacing) appeared, which were attributed to the (100) and (110) crystal planes of δ-MnO2 or birnessite consisting of discrete layers or poorly ordered stacking of adjacent layers, respectively (Brock et al., 1999; Tu et al., 2014). Six reflections in XRD pattern of Fe oxide were identical to those of the lepidocrocite (00–044–1415, PDF number) and similar to other FeOOH standards, demonstrating Fe oxide formed in situ was FeOOH. The BMFO sample only displays three weak reflections (belonged to the BMO part) in the XRD profile, suggesting its poor crystallinity.

### 3.6. XANES and EXAFS analysis

The valence states and coordinating atoms of Fe, Mn, and As in the solid phases of the BFMO, BMO, and Fe oxide were investigated using K-edge XANES and EXAFS analyses. Fig. 6a shows that both the pre-edge peaks and the post-edge regions of Mn K XANES in the BFMO and BMO are almost identical to those of the MnO2 reference samples, with a maximum absorbance at 6561 eV, demonstrating that Mn is primarily in the +IV state in the BFMO and BMO. The adsorption edge energies of Fe species in the BFMO and Fe oxide were between the values of Fe3O4 and Fe2O3, suggesting the mixed valence state of the Fe species due to incomplete Fe(II) oxidation in the BFMO and Fe oxide. Fig. 6c shows that the As K XANES spectra in the As-adsorbed BFMO and BMO are identical to that of Na3AsO4, demonstrating that As(III) adsorbed on the surface of BMO and BFMO was completely oxidized to As(V). However, for the As(III)-Fe oxide sample, the maximum absorbance energy was between those of Na3AsO4 and Na2AsO3, suggesting that only a portion of As(III) was oxidized to As(V) on the Fe oxide surface.

The k²-weighted and Fourier-transformed k³-weighted (R space) Mn, Fe, and As K-edge EXAFS spectra are shown in Fig. S6 and Fig. 7, respectively. Only the first shell (As-O) for the As K-edge spectra and the first and second shells for the Mn and Fe K-edge spectra were fitted due to the small scattering effect of photoelectrons of As coordination (Fig. 7). Good curve-fitting degrees (R factor ≤ 0.012, Tables S2-S4) were obtained between experimental data and theoretical results. The coordination numbers and bond distances are also shown in Tables S2-S4. According to the k²-weighted Mn K-edge spectra (Fig. S6a), the spectrum of BMO is similar to that of MnO2 reference sample. The occurrence of two distinct feature peaks at 8 Å⁻¹ and 9 Å⁻¹ suggests that the BMO structure is a hexagonal birnessite (Butterfield et al., 2013; Webb et al., 2005). The Fe–Fe interatomic distance in Fe-containing samples was approximately 3.06 Å (Table S3), corresponding to edge sharing between Fe³⁺O₆ octahedra (Manceau and Combes, 1988; Thoral et al., 2005). The As-O bond distance of the first shell was found to be 1.68–1.71 Å for As(III&V) adsorbed by the BMO or BFMO, whereas it increased to 1.75 Å for the As(III)-FeOOH sample (Table S4). This finding confirmed the oxidation of As(III) in the BMO or BFMO (Zhang et al., 2014). The similarity of the Fe and Mn K-edge spectra and bond distances of the first and second coordination shells between before and after As(III&V) adsorption suggested that the BFMO, BMO, and FeOOH maintained their local structures during the oxidation or adsorption process. Due to the low concentration of As in the reaction systems, obvious peaks representing the As-O-Fe or As-O-Mn bond in the As K-edge EXAFS spectra were not identified (Fig. 6c and Fig. S6c). However, by comparing the BFMO structure with those of synthetic Fe-Mn oxides (Zhang et al., 2014) and other MnO2 (Manning et al., 2002) and FeOOH samples (Manning et al., 1998; Sherman and Randall, 2003), it can be speculated that the As(III)-
Fig. 3. Oxidation and adsorption of As species in different reaction systems. (a) As(III)+Mn(II)+QJX-1; (b) As(V)+Mn(II)+QJX-1; (c) As(III)+Fe(II)+QJX-1; (d) As(III)+Fe(II)+QJX-1; (e) As(III)+Fe(II)+Mn(II)+QJX-1; (f) As(V)+Fe(II)+Mn(II)+QJX-1. Control: the media containing only QJX-1 and As species. Data points represent the mean value (three replicates) ± standard deviation.
As(V)-adsorbed BFMO samples were inner-sphere bidentate binuclear corner-sharing As-BFMO complexes.

4. Discussion

4.1. BFMO formation

Fe(II) was rapidly oxidized to a hydrous ferric oxide in the aerated environment due to the chemical reaction with oxygen. However, the formation of BMO exhibited a lag time, even after the log phase of bacterial growth (Fig. S1). The inconsistency of bacterial growth and the Mn(II) oxidation rate indicates that QJX-1 exerted an indirect role on Mn(II) oxidation, i.e., superoxide was first formed by the bacteria and then superoxide oxidized Mn(II) to the BMO (Hansel et al., 2012; Learman et al., 2013). Hence, the Fe oxide and BMO parts of the BFMO were formed in turn, and then they were mixed to form the BFMO. It seems that the BMO and Fe oxide parts in BFMO were not chemically bound because their structures were not changed after mixing (Fig. S6). Based on the results of XRD and EXAFS analyses, the BMO formation at circumneutral pH resulted in a highly disordered layered phyllosilicate phase with hexagonal symmetry, which is similar to other BMOs (Bargar et al., 2000, 2009). The Fe oxide part (identified as FeOOH) had octahedral FeO₆ and showed a high affinity for As(III&V) adsorption.

4.2. Mechanism for As(III) oxidation and adsorption by BFMO formed in situ

Based on our results, we obtained a clear picture for the As(III) oxidation and adsorption processes and proposed the following mechanism: As(III) is first adsorbed on the surface of Fe oxide (FeOOH) formed in situ. The FeOOH by itself has a small oxidation ability and some of As(III) ions are oxidized to As(V) on the surface of FeOOH precipitates (Fig. 6). As(III) in the aqueous phase is not oxidized (Fig. 3c). Subsequently, the adsorbed As(III) on the FeOOH surface and the As(III) ions in solution are then oxidized by the formed BMO. The BMO can also oxidize the remaining Fe(II) in solution. The formation of new Fe oxide would further adsorb the As(V) ions remain. Taken together, we conclude that the BMO part in the BFMO is mainly responsible for As(III) oxidation and the FeOOH part dominates As adsorption (Figs. 3 and 6). In comparison with the BMO and FeOOH separately, the formed BFMO had an oxidation and adsorption ability for As(III) that was between the individual parts due to the antagonistic role (occupying oxidation or adsorption sites for each other) between Fe oxide and the BMO.

Previous studies found that BMOs can efficiently adsorb various metal cations (e.g., Pb, Cd, Zn, and Ni) (Nelson et al., 2002; Wang et al., 2009; Zhou et al., 2015). In our study, the formed BMO did not significantly adsorb As (Fig. 2a–b) probably due to that As is an oxyanion. Moreover, we did not observe a shift in As adsorption-desorption during the reaction of the BMO and As species, which has been reported previously in a Mn-oxidizing fungus study (Tani et al., 2004). This discrepancy was probably due to the facts that both the BMO surface and As(V) have negative charges (Fig. 4) and the charge repulsion led to a weak adsorption.

4.3. Comparison between the BFMO and synthetic Fe-Mn oxides

Recently, synthetic Fe-Mn oxides have been demonstrated to remove organics (Liu et al., 2012), inorganics (Peng et al., 2009), and...
metalloids (Zhang et al., 2007) effectively. Here, we compared the characteristics of the BFMO and the synthetic Fe–Mn oxides studied previously (Shan and Tong, 2013; Zhang et al., 2014). The similarities between the two complexes are as follows: (i) they are both formed from a mixture of Mn oxide and Fe oxide precipitation (ii) the optimum Mn/Fe ratio for As adsorption is approximately 1:3 (mol/mol); and (iii) the mechanisms of oxidation and adsorption are identical, i.e., the Mn oxide part controls oxidation and the Fe oxide part dominates adsorption. Due to the differences in the structure of Mn oxide part (the Mn oxide in synthetic Fe–Mn oxides was identified as MnO$_x$, $1.5 < x < 2$ (Zhang et al., 2014)) and the presence/absence of microbes, the BFMO and synthetic Fe–Mn oxides have significant differences in oxidation and adsorption. The dissimilarities are as follows: (i) the BFMO prefers to adsorb As(V) over As(III), but synthetic Fe–Mn oxides have the opposite preference; (ii) synthetic Fe–Mn oxides have a greater adsorption capacity than the BFMO because the BFMO contains Mn-oxidizing bacteria (Mn oxide particles form on the bacterial surface (Jones et al., 2012)), and these bacteria occupy some adsorption sites of FeOOH; (iii) although both the BFMO and synthetic Fe–Mn oxides release Mn(II) during the oxidation of As(III), in the synthetic Fe–Mn oxide system, the released Mn(II) is retained in the water. But in the BFMO system, the released Mn(II) is re-oxidized by the Mn-oxidizing bacteria. With this cycle, the Mn(II) can be eliminated from water. Also due to that BMO could oxidize Fe(II), the Fe(II) would not be retained in water in case the reduction reaction of Fe oxide occurs.

4.4. Implication for Mn, Fe, and As remediation

In natural aquatic ecosystems, Mn oxide often exists with Fe oxide, and they form Fe–Mn oxides together in the sediment (Zhang et al., 2008). Mn-oxidizing microbes play an essential role in the formation of Fe–Mn oxides as microbial Mn oxidation is the dominant pathway in natural Mn oxidation (He et al., 2008). Hence, according to our study, BFMOs may be a critical factor in the natural remediation of As-contaminated water. In addition, high concentrations of Fe(II) and Mn(II) may occur together in As-contaminated groundwater. In conventional groundwater treatment plants, if the presence of Mn-oxidizing microbes, the BFMO could form on the surface of sand filter materials (Bai et al., 2013). The formed BFMO plays a certain role in As removal, and, at the same time, Fe(II) and Mn(II) would not be released. Based on our findings, we propose the development of two strategies to improve groundwater treatment: (i) using bioaugmentation techniques (Bai et al., 2011) (adding some harmless Mn-oxidizing microorganisms to the sand filter); and (ii) using synthetic Fe–Mn oxides as a filter material instead of the sand filter as well as adding harmless Mn-oxidizing microorganisms to the filter in order to prevent the release of Mn(II) and Fe(II) due to the reduction reaction. Compared with current treatment methods, e.g., coagulation/flocculation, iron exchange, adsorption, and chemical redox (Ungureanu et al., 2015), our proposed methods based on BFMO formation would be cost-effective, i.e. low cost with adding bacteria, simultaneous removal of Fe(II), Mn(II) and other pollutants, and without second pollution (BFMOs would be retained in the filter and the released bacteria in case) can be killed by the following UV process.

5. Conclusions

In this study, the proposed hypothesis that the BFMO formed in situ could oxidize and adsorb As species was verified. In addition, the underlying mechanism for As oxidation and adsorption by the BFMO was investigated. There were three main outcomes from the study: (i) BFMO formation occurred through precipitation, i.e., Fe(II) chemical oxidation to FeOOH and Mn(II) biological oxidation to MnO$_2$; the BFMO formed in situ preferred to adsorb As(V) rather than As(III); in general, the optimum initial Mn(II):Fe(II) ratio for As adsorption was approximately 1:3 (mol/mol); (ii) in the BFMO, the BMO part was mainly responsible for As(III) oxidation (a small
Fig. 7. EXAFS spectra of K edge in (a) Mn-containing, (b) Fe-containing, and (c) As-adsorbed BFMO samples and reference samples. Fourier transforms of $\chi(k) k^2$ into the R space, where the red dashed lines denote the curve-fitting results.
contribution from FeOOH), and the FeOOH part dominated As adsorption. It was most likely that the As(II/III/IV)-BFMO complex was an inner-sphere bidentate binuclear corner-sharing (bridged) complex; (ii) As oxidation by synthetic Fe-Mn oxides may release Mn(II) and Fe(II) ions. However, in the BFMO, the released Mn(II) could be re-oxidized by bacteria and the BMO (re)generated in situ could further oxidize As(III) and Fe(II) ions. Therefore, these reactions form a cycle, which can eliminate Fe(II), Mn(II), and As species from the water simultaneously.

Hence, according to our study, the BFMO may be a critical factor in the natural remediation of As-contaminated water as well as in the purification of groundwater for drinking (sand filter process). Furthermore, when groundwater contains high concentrations of Mn(II), Fe(II), and As (III or V), we may use a bioaugmentation technique to eliminate or decrease the above pollutants.

Acknowledgments

We warmly thank Lirong Zheng from the Beijing Synchrotron Radiation Facility for his kind assistance in XAS analysis. This study was supported by a major program (51290282) and a general program (51575837) granted by the National Natural Science Foundation of China, a special fund (13Z03ESPCT) granted by the Key Laboratory of Drinking Water Science and Technology.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.watres.2016.03.068.

References

Adams, L.F., Ghiore, W.C., 1985. Influence of manganese on growth of a sheathless strain of Leptothrix-Discophora. Appl. Environ. Microbiol. 49 (3), 556–562.
Bai, Y.H., Liu, R.P., Liang, J.S., Qu, J.H., 2013. Integrated metagenomic and physicochemical analyses to evaluate the potential role of microbes in the sand filter of a drinking water treatment system. Plos One 8 (4), e61011.
Bai, Y.H., Sun, Q.H., Sun, R.H., Wen, D.H., Tang, X.Y., 2011. Bioaugmentation and adsorption treatment of coking wastewater containing pyridine and quinoline using zeolite-biological aerated filters. Environ. Sci. Technol. 45 (5), 1940–1948.
Bargar, J.R., Fuller, C.C., Marcus, M.A., Brearely, A.J., De la Rosa, M.P., Webb, S.M., Caldwell, W.A., 2009. Structural characterization of terrestrial microbial Mn(II) oxide and As(V) Geochim. Cosmochim. Ac 73 (4), 889–910.
Bargar, J.R., Tebo, B.M., Vilinski, J.E., 2000. In situ characterization of Mn(II) oxidation by spores of the marine Bacillus str. sp. strain GB-1. Geochim. Cosmochim. Ac 64 (16), 2775–2778.
Berg, M., Li, J., Torgersen, P.T.K., Trang, P.T.K., Viet, P.H., Giger, W., Stuben, D., 2006. Arsenic removal from groundwater by bacterial Mn-oxidizing fungus, strain KR21-2. Environ. Sci. Technol. 38 (24), 6618–6624.
Bolos, D., Biser, J., Maradiaga, J., Fonda, E., Nahon, D., Bottero, J.Y., 2005. XAS study of arsine(III) and arsenic(III) oxidation by magnetic nanoparticles modified with Fe-Mn binary oxide. Water Res. 47 (10), 3411–3421.
Sherman, D.M., Randall, S.R., 2003. Surface complexation of arsenic(III) on hydroxide(II) and iron(III) in dilute acid solution by ozone. Hydrometallurgy 62 (1–2), 51–52.
Bai, L., Ruan, J.H., Xiao, R., Gao, H.F., Liu, F.P., 2012. Removal of tetracycline from water by Fe-Mn binary oxide. J. Environ. Sci. China 24 (2), 242–247.
Manceau, A., Combes, J.M., 1988. Structure of Mn and Fe oxides and oxyhydroxides: a topological approach by EXAFS. Phys. Chem. Miner. 15 (3), 283–295.
Manning, B.A, Fendorf, S.E., Bostick, B., Suarez, D.L., 2002. Arsenic(III) oxidation and (V) adsorption reactions on synthetic birnessite. Environ. Sci. Technol. 36 (5), 976–981.
Manning, B.A, Fendorf, S.E., Goldberg, S., 1998. Surface structures and stability of arsenic(III) on goethite: spectroscopic evidence for inner-sphere complexes. Environ. Sci. Technol. 32 (16), 2383–2388.
Murray, K.J., Mozzafarzadeh, M.L., Tebo, B.M., 2005. Cr(III) oxidation and Cr toxicity in cultures of the manganese(II)-Oxidizing Pseudomonas putida strain GB-1. Geo- microb. J. 22 (3–4), 151–159.
Nelson, Y.M., Lion, L.W., Shuler, M.L., Ghiore, W.C., 2002. Effect of oxide formation mechanisms on lead adsorption by biogenic manganese (hydr)oxides, iron (hydr)oxides, and their mixtures. Environ. Sci. Technol. 36 (3), 421–425.
Newville, M., 2001. FEFFIT: interactive EXAFS analysis and FEFF fitting. J. Synchrotron Rad. 8, 322–324.
Nishimura, T., Umetu, N., 2001. Oxidative precipitation of arsenic(III) with man- ganese(II) and iron(II) in dilute acid solution by ozone. Hydrometallurgy 62 (1–2), 51–52.
Postawa, A., Hayes, C., 2013. Best Practice Oguide on the Control of Iron and Manganese in Water Supply. IWA Publishing, London, UK.
Shan, C., Tong, M.P., 2013. Efficient removal of trace arsenite through oxidation and adsorption by magnetic nanoparticles modified with Fe-Mn binary oxide. Water Res. 47 (10), 3411–3421.
Tian, Y., Miyata, N., Ohashi, T., Sayauma, H., Iwahori, K., Soma, M., 2004. Interaction of inorganic arsenic with biogenic manganese oxide produced by Mn(III/II)-oxidizing bacteria, strain K921-2. Environ. Sci. Technol. 38 (24), 6618–6624.
Tebo, B.M., Spiro, T.G., Bargar, J.R., Spoto, G., 2010. Bacteriogenic manganese oxides. Accounts Chem. Res. 43 (1), 2–9.
Thoral, S., Rose, J., Garnier, M.J., Van Geen, A., Refait, P., Traverse, A., Fonda, E., Nahon, D., Bottero, J.Y., 2005. XAS study of iron and arsenic speciation during Fe(II) oxidation in the presence of As(III). Environ. Sci. Technol. 39 (24), 9478–9485.
Tani, Y., Miyata, N., Ohashi, T., Sayauma, H., Iwahori, K., Soma, M., 2004. Interaction of inorganic arsenic with biogenic manganese oxide produced by Mn(III/II)-oxidizing bacteria, strain K921-2. Environ. Sci. Technol. 38 (24), 6618–6624.
Voegelin, A., Kaegi, R., Berg, M., Nitzsche, K.S., Anulódo, J.A., 1999. Arsenic in drinking water problems and solutions. Water Sci. Technol. 40 (2), 69–76.
Vira-raghavan, T., Subramanian, K., Anulódó, J.A., 1999. Arsenic in drinking water problems and solutions. Water Sci. Technol. 40 (2), 69–76.
Zhang, G.S., Liu, F.D., Liu, H.J., Qu, J.H., Liu, R.P., 2014. Respective role of Fe and Mn oxide contents for arsenic sorption in iron and manganese binary oxide: an X-ray absorption spectroscopy investigation. Environ. Sci. Technol. 48 (17), 10316–10322.
Zhang, G.S., Ko, S.O., 2015. Heavy metal adsorption with biogenic manganese oxides generated by Pseudomonas putida strain MnB1. J. Ind. Eng. Chem. 24, 132–139.

Y. Bai et al. / Water Research 98 (2016) 119–127