Repeated Anaerobic Microbial Redox Cycling of Iron

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Received 7 February 2011/ Accepted 28 June 2011

Some nitrate- and Fe(III)-reducing microorganisms are capable of oxidizing Fe(II) with nitrate as the electron acceptor. This enzymatic pathway may facilitate the development of anaerobic microbial communities that take advantage of the energy available during Fe-N redox oscillations. We examined this phenomenon in synthetic Fe(III) oxide (nanocrystalline goethite) suspensions inoculated with microflora from freshwater river floodplain sediments. Nitrate and acetate were added at alternate intervals in order to induce repeated cycles of microbial Fe(III) reduction and nitrate-dependent Fe(II) oxidation. Addition of nitrate to reduced, acetate-depleted suspensions resulted in rapid Fe(II) oxidation and accumulation of ammonium. High-resolution transmission electron microscopic analysis of material from Fe redox cycling reactors showed amorphous coatings on the goethite nanocrystals that were not observed in reactors operated under strictly nitrate- or Fe(III)-reducing conditions. Microbial communities associated with N and Fe redox metabolism were assessed using a combination of most-probable-number enumerations and 16S rRNA gene analysis. The nitrate-reducing and Fe(III)-reducing cultures were dominated by denitrifying Betaproteobacteria (e.g., Dechloromonas) and Fe(III)-reducing Deltaproteobacteria (Geobacter), respectively; these same taxa were dominant in the Fe redox cycling cultures. The combined chemical and microbiological data suggest that both Geobacter and various Betaproteobacteria participated in nitrate-dependent Fe(II) oxidation in the cycling cultures. Microbially driven Fe-N redox cycling may have important consequences for both the fate of N and the abundance and reactivity of Fe(III) oxides in sediments.

Oxygen, nitrate, and Mn(IV) oxides are the primary oxidants for Fe(II) in natural systems. Fe(II) is subject to spontaneous chemical oxidation by oxygen and Mn(IV) oxides at circumneutral pH (16, 31, 33, 36). In contrast, the abiotic reaction of Fe(II) with nitrate is negligible under the temperature and aqueous geochemical conditions typical of natural soil and sedimentary environments (58). Although the presence of nitrate-reducing, Fe(II)-oxidizing organisms has been documented in a wide variety of environments (5, 13, 25, 28, 37, 40, 49, 51, 52), little is known about the quantitative coupling of Fe and N redox cycles in sediments. In particular, virtually nothing is known about Fe-N redox cycling in sediments subject to spatial or temporal shifts in redox conditions.

The input of oxidants compared to utilizable organic carbon controls the predominant terminal electron-accepting pathway (TEAP) in a given soil or sediment horizon. Where the fluxes of oxidants and organic carbon are relatively constant, sequential consumption of electron acceptors leads to the formation of stable redox gradients (3, 20). Many subsurface environments that are very active hydrologically and which support a diverse microflora (11), e.g., local-flow groundwater systems (12), however, are characterized by large fluctuations in chemical conditions. Redox fluctuations in such environments are likely to have a critical impact on Fe species content and thereby strongly influence the behavior of natural and contaminant compounds whose behavior is linked to Fe redox cycling.

Nitrate-dependent Fe(II) oxidation enhances the potential for coupled Fe-N redox interactions in sediments and may be associated with the development of microbial populations specifically adapted to take advantage of the energy available during redox oscillations (53, 60). Transient redox fluctuations can also have a significant impact on Fe(III) oxide mineralogy (15), although the cumulative effects of such fluctuations are not well known (54). In this paper we present the results of batch experiments designed to explore geochemical and microbiological phenomena associated with anaerobic Fe-N redox cycling in reaction systems experiencing repeated fluctuations in organic carbon (acetate) and nitrate loading.

MATERIALS AND METHODS

Culture medium. O2-free, N2-bubbled, 1,4-piperazinediethanesulfonic acid (PIPES)-buffered (10 mM, pH 6.8) artificial groundwater (AGW) medium (60) containing (per liter) 5.5 g (62 mmol) goethite (α-FeOOH) synthesized by air oxidation of FeCl2·2H2O in NaHCO3 buffer (44) was used in all experiments. This goethite corresponds to the “high-surface goethite” (surface area, ca. 150 m2 g−1; average particle size, 15 to 30 nm) discussed in reference 43. Previous studies have demonstrated that the abundance of amorphous impurities (e.g., ferrihydrite) is very small in this material (42), and transmission electron microscopy (TEM) analysis verified the absence of amorphous phases in the starting material (see below). Goethite rather than ferrihydrite was used in these experiments because ferrihydrite typically undergoes major phase transformations during microbial reduction (61), which could have complicated assessment of the potential for nitrate-driven Fe redox cycling.

Incubation experiments. Three experiments were conducted in duplicate. System 1 (nitrate reducing) was amended with limiting acetate (2.0 mM) relative to nitrate (2.2 mM). System 2 (Fe(III) reducing) was amended with 2.0 mM acetate only. System 3 (Fe redox cycling) was initially amended with 2.1 mM acetate and 1.0 mM nitrate, and upon depletion, subsequent additions of either acetate or nitrate were made to induce Fe redox cycling. The system 1 and system 2 incubations lasted for only a few weeks, as nitrate reduction and Fe(III) reduc-
tion ceased after ca. 1 and 2 weeks, respectively. In contrast, the Fe redox cycling experiment went on much longer (140 days) in order to assess the potential for repeated coupled Fe/N redox metabolism.

Acetate and nitrate were added from sterile, anaerobic stock solutions. Each reactor was inoculated with 5 ml of fresh surface sediment (0 to 2 cm) collected from near the banks of the Wisconsin River, three miles west of Portage, WI (close to site S3 described by Forsfjord and Stanley [19]). The sediment contained approximately equal concentrations (ca. 20 mmol liter⁻¹) of 0.5 M HCl-extractable Fe(II) and Fe(III), indicating active Fe(III) reduction (30). Although sediment pore fluids were depleted in nitrate, the section of the Wisconsin River from which the inoculum was obtained contains ca. 0.1 mM nitrate, and floodplain sediments are strong sinks for nitrate (19).

Samples for determination of Fe(II) (aqueous and solid associated), nitrate, ammonium, and acetate analyses were taken immediately after bottles were inoculated and then every 2 to 5 days for up to 135 days. Between sampling periods, the bottles were stored in the dark at room temperature without shaking. The contents of the bottles were resuspended before taking 1.5-ml samples with an N₂-flushed syringe. The aqueous phase was separated by centrifugation (12,000 × g for 10 min) in an anaerobic chamber (95% N₂, 5% H₂). For microbial community analysis, 5-ml samples were removed at the beginnings and ends of the experiments and stored anaerobically at –20°C.

Enumeration, enrichment, and isolation. At the ends of the incubation experiments, bacterial populations were enumerated as CFU using an anaerobic roll-tube method (26). Ten-fold serial dilutions were made in anaerobic AGW; 1-ml subsamples from each dilution were introduced into duplicate tubes containing 7 ml of sterile, anaerobic AGW liquefied agar (1.5% at 50°C) with the appropriate electron donor/acceptor. Enumerations were made for (i) nitrate-reducing (5 mM nitrate, 10 mM acetate), (ii) autotrophic nitrate-reducing Fe(II)-oxidizing (0.5 mM acetate, 5 mM nitrate, 5 mM FeCl₂, 2H₂O) [referred to here as Fe(II)], (iii) autotrophic nitrate-reducing, Fe(II)-oxidizing [5 mM nitrate, 5 mM FeCl₂ (II)], and (iv) Fe(III)-reducing [80 mmol liter⁻¹ synthetic amorphous Fe(III) oxide (29), 10 mM acetate] organisms. The contents of the inoculated tubes were mixed, and the tubes were spun rapidly on a Bellco tube spinner until the agar had cooled on the sides of the culture tube. After 4 to 5 weeks of incubation in the dark at room temperature, colonies formed on the sides of the tubes were enumerated and the CFU per ml of inoculum were calculated.

Acetate-utilizing nitrate- and Fe(III)-reducing bacteria were enriched in AGW medium with 5 mM acetate, 50 mg liter⁻¹ yeast extract (YE), and either 5 mM nitrate or 100 mM sodium thiosulfate per liter. Nitrate reducers were isolated following sequential transfers of anaerobic enrichment cultures on AGW medium with 5 mM nitrate and 5 mM acetate. Single colonies from a highly enriched culture were selected from medium solidified with 1.5% Noble agar. Nitrate-reducing isolates were identified phylogenetically based on their 16S rRNA gene sequences as described below.

Stable Fe(II)-reducing cultures were obtained from all three systems. Roll-tube isolation procedures with either (i) acetate and amorphous Fe(III) oxide or (ii) a combined mixture (referred to as “nfa mix”) of electron donors and acceptors (5 mM pyruvate, 10 mM acetate, 5 mM nitrate, 10 mM fumarate, 25 mg liter⁻¹ YE) were applied to test isolation of dissimilatory iron-reducing bacteria (DIRB) for each system. Pure cultures so obtained were grown in nfa medium with 5 mM acetate, 50 mg liter⁻¹ yeast extract (YE), and either 5 mM nitrate or 100 mM sodium thiosulfate per liter. Nitrate reducers were isolated following sequential transfers of anaerobic enrichment cultures on AGW medium with 5 mM nitrate and 5 mM acetate. Single colonies from a highly enriched culture were selected from medium solidified with 1.5% Noble agar. Nitrate-reducing isolates were identified phylogenetically based on their 16S rRNA gene sequences as described below.

16S rRNA gene analysis. Subsamples removed from duplicate reactors were pooled prior to extraction of genomic DNA using the MoBio UltraClean soil DNA kit. Materials from the initial (time zero) samples were similarly pooled and extracted. Purified DNA was amplified using the universal bacterial primer set GMSF-GMAR (35), resulting in a PCR product length of approximately 1.5 kb. DNA extracts from acetate/goethite enrichment cultures were amplified using 8F and 907R primers. PCR products were purified using a MinElute PCR purification kit. The purified product was ligated using a Promega pGEM-T Easy Vector System kit, transformed into competent Escherichia coli cells, and plated on LB agar containing 100 mg ml⁻¹ ampicillin and X-Gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside). Transformants were incubated at 37°C overnight, and colonies with recombinant vectors were selected. Recombinant vectors were grown overnight in LB broth at 37°C. Plasmids were isolated using a Promega Wizard DNA purification kit and sequenced at the University of Wisconsin Biotechnology Center.

DNA extracts from several samples were sent to Microbial Insights, Inc., for denaturing gradient gel electrophoresis (DGGE) analysis of 16S rRNA genes as previously described (10). Prominent bands were excised, and the DNA contained in them was reamplified and sequenced.

Sequence data from the clone library and DGGE analyses were manually edited using the BioEdit program (version 7.0.5.3) (24) to remove vector sequences and eliminate poorly resolved sequences. Whenever possible, edited sequences were assigned genus-level taxonomic affiliation by way of BLAST (1) searches. A value of 95% similarity was used as a cutoff for assignment of genus-level affiliation (21). In a few cases, only family-level affiliations could be assigned. In one instance, a collection of five diverse Alphaproteobacteria from the Fe(III)-reducing cultures were grouped together as “other Alphaproteobacteria.”

Analytical methods. Acetate, nitrate, and nitrite concentrations were measured using a Dionex 1000 ion chromatograph. Ammonium was quantified colorimetrically (46). Aqueous Fe(II) was measured by adding an aliquot of supernatant directly to ferrozine (48) in the anaerobic chamber. Sorbed Fe(II) was measured by decanting the remaining supernatant and replacing it with an equal volume of 0.5 M HCl; the pellet was resuspended and placed on a shaker table at 100 rpm for 1 h, and Fe(II) in the extract was measured (after centrifugation) using ferrozine.

TEM. Fe(III) oxides from the batch reactors were recovered by centrifugation at 10,000 × g for 10 min and washed with anoxic deionized water. Samples were placed on copper grids, covered with a carbon-coated Formvar film, and examined with a Philips CM 200UT transmission electron microscope (spherical aberration coefficient, 0.5 mm; point-to-point resolution, 0.19 nm) equipped with Noran Voyager X-ray energy-dispersive spectroscopy (EDS).

RESULTS AND DISCUSSION

Nitrate-reducing cultures. Acetate and nitrate were consumed concomitantly in the acetate-limited, nitrate-reducing incubations (system 1) (Fig. 1A). Acetate was depleted by day 6, whereas some nitrate remained for the duration of the experiment. No increase in Fe(II) was observed, and the concentration remained similar to the background level of ca. 0.2 mM. These results indicate that denitrification was the dominant pathway for acetate metabolism in these cultures. This conclusion is consistent with the microbial community analysis data (Fig. 2B; see Fig. S1 in the supplemental material), which...
showed a predominance of phylotypes related to known Beta-proteobacteria denitrifying taxa (e.g., Dechloromonas). After 18 days of incubation, the nitrate-reducing cultures contained substantial numbers of both acetate-oxidizing nitrate reducers and nitrate-dependent Fe(II) oxidizers that could grow with or without acetate (Table 1); lower but significant numbers of Fe(III)-reducing organisms were also detected. Four nitrate-reducing Betaproteobacteria were isolated from the acetate/nitrate cultures (see Table S1 in the supplemental material), each of which reduced nitrate without ammonium production when grown with 5 mM acetate and 5 mM nitrate. One of the cultures (UWNR4, a Dechloromonas strain) was able oxidize a significant amount (ca. 90%) of Fe(II) in mixotrophic medium containing 5 mM nitrate, 5 mM Fe(II), and 0.5 mM acetate (data not shown).

**Fe(III)-reducing cultures.** Approximately 6 mmol of HCl-extractable Fe(II) per liter was produced during metabolism of 1 mM acetate in the nitrate-free, Fe(III)-reducing (system 2) cultures (Fig. 1B). These cultures were dominated by Geobac-

![Diagram of 16S rRNA gene clone libraries from various cultures](image)

**FIG. 2.** Compositions of 16S rRNA gene clone libraries from the original sediment inoculum (95 clones) (A), the nitrate-reducing cultures after 11 days of incubation (112 clones) (B), the Fe(III)-reducing cultures after 18 days of incubation (98 clones) (C), and the Fe cycling cultures after 129 days of incubation (100 clones) (D). Numbers in parentheses indicate the number of clones for each taxonomic assignment and their percent abundance in the library.

**TABLE 1.** Roll tube enumeration results for the nitrate-reducing, Fe(III)-reducing, and Fe cycling cultures

<table>
<thead>
<tr>
<th>Culture system</th>
<th>Electron donor, acceptor</th>
<th>Cell density (cells ml⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (nitrate reducing)</td>
<td>Acetate, nitrate</td>
<td>3.4 × 10⁷</td>
</tr>
<tr>
<td></td>
<td>Acetate, Fe(III)</td>
<td>6.0 × 10⁵</td>
</tr>
<tr>
<td></td>
<td>Fe(II), nitrate</td>
<td>2.1 × 10⁷</td>
</tr>
<tr>
<td></td>
<td>Fe(II)/acetate, nitrate</td>
<td>1.5 × 10⁸</td>
</tr>
<tr>
<td>2 [Fe(III) reducing]</td>
<td>Acetate, nitrate</td>
<td>ND³</td>
</tr>
<tr>
<td></td>
<td>Acetate, Fe(III)</td>
<td>1.0 × 10⁶</td>
</tr>
<tr>
<td></td>
<td>Fe(II), nitrate</td>
<td>5.2 × 10⁵</td>
</tr>
<tr>
<td></td>
<td>Fe(II)/acetate, nitrate</td>
<td>4.0 × 10⁵</td>
</tr>
<tr>
<td>3 (Fe cycling)</td>
<td>Acetate, nitrate</td>
<td>5.3 × 10⁶</td>
</tr>
<tr>
<td></td>
<td>Acetate, Fe(III)</td>
<td>1.5 × 10⁷</td>
</tr>
<tr>
<td></td>
<td>Fe(II), nitrate</td>
<td>7.2 × 10⁶</td>
</tr>
<tr>
<td></td>
<td>Fe(II)/acetate, nitrate</td>
<td>6.3 × 10⁶</td>
</tr>
</tbody>
</table>

³ All cultures were incubated for 2 months prior to colony enumeration. Results are means for duplicate tubes; the ranges for duplicates were <5% of the means.

³ ND, not determined.
second nitrate addition at day 36 facilitated continued nitrate-dependent Fe(II) oxidation. Subsequent addition of acetate at day 50 led to rapid nitrate depletion followed by another pulse of Fe(III) reduction. Addition of nitrate at day 74 under acetate-depleted conditions led to rapid Fe(II) oxidation. Acetate added again 4 days later resulted in immediate regeneration of the Fe(II). Another nearly identical cycle of nitrate-driven Fe(II) oxidation and acetate-fueled Fe(III) reduction ensued, followed by a final round of Fe(II) oxidation. A total of four complete cycles of Fe reduction and oxidation were thus obtained in this experiment. The time needed to complete an Fe redox cycle decreased from about 50 days for the first cycle to 20 to 25 days in subsequent cycles, suggesting that Fe redox cycling populations were stimulated. Nitrate-dependent Fe(II) oxidation was associated with ammonium accumulation (Fig. 3D). Nitrite was detected only sporadically and always at concentrations less than 50 μM, which suggests that nitrate was in general reduced directly to ammonium (or other reduced end products, e.g., N₂; see below).

The 4-month-old Fe cycling cultures yielded 15- to 20-fold more DIRB than the nitrate-reducing and Fe(III)-reducing cultures operated for a much shorter period (Table 1), which is consistent with the much larger total amount of Fe(III) oxide reduction activity that took place in the cycling reactors. The initial microbial community in the Fe cycling cultures closely resembled that in the nitrate-reducing cultures (see Fig. S1B and D in the supplemental material). DGGE revealed the development of a more diverse community during Fe redox cycling (see Fig. S1D to H in the supplemental material), which was confirmed by examination of the clone library constructed at the end of the experiment (Fig. 2D).

**Mechanisms of anaerobic Fe redox cycling.** Fe(II)-driven nitrate reduction produced ammonium, as observed in a previous analogous study (60). The overall molar ratio of ammonium produced to Fe(II) oxidized (0.17 ± 0.01) (Fig. 3D, inset) was similar to that expected for the oxidation of aqueous (Fe²⁺) or surface-associated ([≡O-Fe⁺]) Fe(II) coupled to reduction of nitrate to ammonium according to reactions such as Fe²⁺ + 0.125NO₃⁻ + 1.625H₂O → FeOOH + 0.125NH₄⁺ + 1.75H⁺ and ≡O-Fe⁺ + 0.125NO₃⁻ + 1.625H₂O → ≡OH + FeOOH + 0.125NH₄⁺ + 0.75H⁺. This result suggested involvement of Fe(III)-reducing *Geobacter* species that are capable of oxidizing Fe(II) with reduction of nitrate to ammonium (60).

Extensive, but unsuccessful, attempts were made to recover pure cultures of *Geobacter* from acetate/goethite enrichments derived from the Fe redox cycling reactors in order to determine whether such organisms were capable of Fe(II) oxidation with reduction of nitrate to ammonium. Organisms from the *Rhodocyclaceae* (Azospira and Dechloromonas), the *Desulfovibrionaceae* (Desulfovibrio), and the *Sporomusa-Pectinatus-Selenomonas* phyltic group (*Anaerovibrio*) were recovered from colonies picked from roll tubes (see Table S1 in the supplemental material). None of these isolates grew via Fe(III) reduction [10 mM Fe(III)-NTA or 50 mmol liter⁻¹ of amorphous Fe(III) oxide] or oxidized significant amounts of Fe(II) in medium containing 5 mM nitrate, 5 mM Fe(II), and 0.5 mM acetate.

Despite the lack of recovered acetate-oxidizing DIRB, 16S rRNA gene clone libraries (ca. 800-bp fragments) showed that
the three goethite-reducing enrichment cultures (25 clones each) were dominated by an operational taxonomic unit (OTU) 95% similar to *Geobacter pelophilus* (50). These sequences were 99% similar to several *Geobacter*-related OTUs recovered in 16S rRNA gene libraries from the Fe redox cycling cultures. The three acetate/goethite enrichment cultures were tested for the ability to oxidize Fe(II) with nitrate following an initial period of acetate-limited Fe(III) reduction. These experiments were carried out with cells grown on Fe-rich smectite, which proved to be the best substrate for growth of the later, less robust generations of the enrichments. Cells for these experiments were obtained by transferring the goethite-reducing enrichments to a suspension of specimen smectite (NaU-2) in standard culture medium and growing them for three generations prior to addition of nitrate to acetate-depleted cultures. All three cultures showed significant Fe(II) oxidation activity [20 to 35% of the available Fe(II)] in the presence of excess nitrate. These results support the inference that *Geobacter*-related DIRB participated in nitrate-dependent Fe(II) oxidation in the Fe redox cycling cultures.

Organisms from the Betaproteobacteria known to be capable of nitrate-dependent Fe(II) oxidation (e.g., *Dechloromonas, Dechlorosoma,* and *Thiobacillus* [59]) were also detected in the Fe redox cycling reactors (Fig. 2D; see Fig. S1 in the supplemental material). The quantitative role of these taxa in Fe(II) oxidation is uncertain, given that all known betaproteobacterial nitrate-dependent Fe(II) oxidizers produce N₂ rather than ammonium as the end product of nitrate reduction (59). It is important to note, however, that the ratio of nitrate consumed to Fe(II) oxidized during the last three periods of Fe(II) oxidation (0.25 ± 0.02; see Fig. S2A in the supplemental material), when no acetate was present, was twice that expected for Fe(II) oxidation coupled to reduction of nitrate to ammonium. This result suggests that some of the nitrate consumed during these periods was linked to organotrophic metabolism, e.g., endogenous respiration and/or oxidation of dead bacterial biomass. This conclusion raises the possibility that some of the ammonium accumulation during nitrate-dependent Fe(II) oxidation was linked to organotrophy (e.g., by nitrate-reducing *Geobacter*) rather than Fe(II) oxidation, which in turn suggests that some of the nitrate-driven Fe(II) oxidation activity could have been carried out by betaproteobacterial denitrifiers as opposed to *Geobacter*, including both heterotrophic (e.g., *Dechloromonas* and *Dechlorosoma*) and autotrophic (e.g., *Thiobacillus*) taxa.

The Solver tool in Microsoft Excel was used to constrain the possible quantitative significance of these different pathways for nitrate reduction. The ratio of nitrate consumed to Fe(II) oxidized during the last three periods of Fe(II) oxidation was fixed at the observed value of 0.253 (see Fig. S2A in the supplemental material). The presumed rate of organotrophic nitrate reduction, along with the fraction of nitrate reduced to N₂ versus ammonium during organotrophic and lithotrophic [Fe(II)-driven] nitrate reduction, was then varied within the Solver routine to reproduce the observed ratio of ammonium produced to Fe(II) oxidized during the last three periods of Fe(II) oxidation (0.182; see Fig. S2B in the supplemental material). A summary of the calculation procedure and results is provided in Table S2 in the supplemental material. The results indicated a rate of organotrophic nitrate reduction equivalent to 46% of total nitrate consumption, with 68% of organotrophic nitrate reduction and 83% of Fe(II)-driven nitrate reduction resulting in ammonium production. The percentage of Fe(II) oxidation coupled to reduction of nitrate to ammonium was 76% of the total amount. These calculations lend conceptual support to the hypothesis suggested by the DGGE and clone library results, i.e., that nitrate-driven Fe(II) oxidation was carried out by both betaproteobacterial denitrifiers and ammonium-producing *Geobacter* spp. in the Fe cycling reactors.

**Influence of Fe redox cycling on Fe(III) oxide mineralogy.**

High-resolution TEM analysis demonstrated the presence of goethite nanocrystals less than 30 nm in length and 10 nm in diameter (Fig. 4). Selected area electron diffraction patterns indicated the relatively low crystallinity of the goethite and the absence of major mineral-phase transformations during reduction or Fe redox cycling (see Fig. S3 in the supplemental material). These results are consistent with a prior study (60) which showed that nitrate-dependent oxidation of microbially reduced goethite did not result in the formation of new crystalline Fe(III) oxide phases. However, TEM images of samples from the Fe redox cycling cultures did show ubiquitous amorphous material surrounding the goethite nanocrystals that were absent in the nitrate- and Fe(III)-reducing systems (Fig. 4). To our knowledge this is the first demonstration of the generation of amorphous Fe(III) oxide coatings on otherwise crystalline Fe(III) oxide mineral phases during nitrate-dependent Fe(II) oxidation. These results are consistent with the view that Fe redox cycling is likely to result in the formation of highly reactive, amorphous Fe(III) oxide phases (55), and they

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**FIG. 4.** High-resolution TEM images of samples collected at the ends of the nitrate-reducing (A), Fe(III)-reducing (B), and Fe redox cycling (C) experiments. Note the marked increase in the abundance of amorphous material surrounding and among goethite nanocrystals in the sample from the Fe redox cycling reactor (C).
may explain the observed production of small quantities of dilute HCl-extractable (presumably amorphous) Fe(III) oxide during nitrate-dependent oxidation of microbially reduced goethite by sediment microflora and Geobacter metallireducens (60).

Environmental implications. Our experiments show that a microbial system undergoing repeated temporal oscillations in the input of organic carbon and nitrate developed a community structure capable of mediating sustained anaerobic Fe redox cycling through dissimilatory Fe(III) oxide reduction and nitrate-dependent Fe(II) oxidation. These results are analogous to those of a previous experiment with natural-sediment microflora in which Fe went through a single cycle of Fe reduction and oxidation (60) and to those of experiments with cocultures of Geobacter bremensis (53) or Geobacter sulfurreducens (7) (neither of which reduce nitrate) and the lithoautotrophic nitrate-reducing, Fe(II)-oxidizing enrichment culture of Straub et al. (52). Together these studies suggest that microbial Fe cycling communities are likely to be present and active in anoxic soil and sedimentary environments experiencing shifts in organic carbon and nitrate input, analogous to those known to be present in aerobic/anoxic interfacial environments (6, 8, 23, 47). When inputs of organic carbon are high compared to nitrate, organic carbon oxidation by nitrate-reducing bacteria exhausts available nitrate, thus allowing microbial Fe(III) reduction to become the predominant TEAP (4, 32, 34, 57). During subsequent periods of reduced organic carbon loading, rates of nitrate respire may exceed rates of organotrophic nitrate reduction, resulting in the potential for lithotrophic, nitrate-dependent Fe(II) oxidation, e.g., at the redox boundary in nitrate-contaminated aquifers in agricultural areas (17, 18, 39) or at the fringe of the Fe(III) reduction zone in organic-contaminated aquifers (2, 14, 56). A similar coupling of N and Fe redox may take place in fresh water or marine surface sediments (27, 51). In each of these situations, reducing equivalents stored in the form of Fe(II)-bearing solid phases (including, in some cases, iron sulfide minerals [9, 22]) may eventually serve as a significant source of energy for lithotrophic microbial metabolism during periods of reduced organic carbon input or during bioturbation- or wave-induced mixing of aquatic surface sediments. The importance of solid-associated Fe(II) in Fe redox cycling is illustrated by the fact that aqueous Fe(II) accounted for less than 20% of total Fe(II) production and oxidation in the Fe(III)-reducing and Fe cycling cultures (Fig. 1B and 3C).

A key aspect of this study and the previous related study (60) is that ammonium was a major end product of nitrate-dependent Fe(II) oxidation during anaerobic Fe redox cycling. This finding has important implications for the fate of N in sediments, as ammonium so produced may eventually be returned to surface waters where it could serve as nutrient for algal and higher plant production. In contrast, conversion of nitrate to N2 effectively represents a loss of N from the sedimentary N cycle (45). Recent studies suggest that this dissipative reduction of nitrate to ammonium may also occur during sulfide-linked nitrate reduction in freshwater wetland sediments (9, 38). Finally, given the fundamental influence of Fe(III) oxide surface area and crystallinity on rates of microbial Fe(III) reduction (41), changes in Fe(III) oxide mineralogy (e.g., production of amorphous surface coatings) associated with microbially Fe redox cycling have the potential to alter the reactivity of Fe(III) oxides in soils and sediments.

ACKNOWLEDGMENTS

We thank Marco Blöthe for his help with 16S rRNA gene sequence analyses and Jason Flagg (Indiana University) for assistance with microbial isolations.

This research was supported by grant 0525510 from the NSF Biogeochemistry Program and grants DE-FG02-06ER64184 and ER64172-1027487-001191 from the Office of Biological and Environmental Research, U.S. Department of Energy, Office of Science.

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