Characteristics and Kinetic Analysis of AQS Transformation and Microbial Goethite Reduction: Insight into “Redox mediator-Microbe-Iron oxide” Interaction Process

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The characteristics and kinetics of redox transformation of a redox mediator, anthraquinone-2-sulfonate (AQS), during microbial goethite reduction by Shewanella decolorationis S12, a dissimilatory iron reduction bacterium (DIRB), were investigated to provide insights into “redox mediator-iron oxide” interaction in the presence of DIRB. Two pre-incubation reaction systems of the “strain S12-goethite” and the “strain S12-AQS” were used to investigate the dynamics of goethite reduction and AQS redox transformation. Results show that the concentrations of goethite and redox mediator, and the inoculation cell density all affect the characteristics of microbial goethite reduction, kinetic transformation between oxidized and reduced species of the redox mediator. Both abiotic and biotic reactions and their coupling regulate the kinetic process for “Quinone-Iron” interaction in the presence of DIRB. Our results provide some new insights into the characteristics and mechanisms of interaction among “quinone-DIRB- goethite” under biotic/abiotic driven.

The high abundance and the redox sensitivity of Fe(III) oxides in natural environment have led to the extensive research in their redox biogeochemical behaviors. The anaerobic environments such as sediments, soils, sediment-water interface and aquifers commonly contain organic matters with sensitive chemical structures that can act as redox mediator. Redox mediators, such as humic substances containing quinone structures, which have the redox potentials to facilitate electron acceptance and donation, can mediate the extracellular electron transferring from a bacterium to an insoluble electron acceptor, such as iron oxides. Extensive researches have consequently been performed to investigate the role of redox mediators in enhancing the degree and extent of microbial iron reduction.

Quinone, as a redox mediator, can be reduced to a corresponding hydroquinone by dissimilatory iron reduction bacterium (DIRB). The hydroquinone can be abiotically oxidized by transferring electrons to electron acceptors, such as Fe(III) oxides including ferrihydrite, lepidocrocite, hematite, and goethite. These previous studies generated good knowledge of how redox mediators such as quinone interact with Fe(III) oxides. Most of these studies, however, focused on the abiotic redox reaction between the reduced form of a redox mediator and iron oxides. These studies have established abiotic kinetics to describe the redox transformation between redox mediators and iron oxides. The kinetic knowledge and the established models have been conceptualized to describe microbial reduction of iron oxides mediated by redox mediators.

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However, research on the coupling biotic and abiotic interactions between reduced redox mediators and iron oxides in the presence of DIRB should be needed further investigation. The reduced redox mediator can be re-generated in the presence of DIRB by reducing oxidized redox mediator to its corresponding reduced form, thus enhancing the reduction capacity and reaction duration. In addition, iron oxides can be directly reduced by DIRB that may affect abiotic reduction by the redox mediator. For example, the Fe(II) produced by the microbial reduction can be adsorbed on residual iron oxide surfaces, which can impact subsequent redox cycle of quinone and iron reduction. Therefore, it is important to understand the behavior of “Quinone-Iron” interaction system in the presence of DIRB.

In our current study, the humic substance containing quinoid structure, anthraquinone-2-sulfonate (AQS), and goethite were selected as the model redox mediator and iron oxide respectively, with the aim at the investigation of the transformation between oxidized and reduced species of redox mediator during the process of microbial reduction of iron oxide. Two pre-incubation reaction systems termed subsequently as the “DIRB-iron oxide” and “DIRB-redox mediator”, were specifically selected to investigate the interactions among “quinone-DIRB-iron oxide” under biotic/abiotic driven. The temporal variations in concentrations of quinone and different species of reductively produced ferrous iron were monitored to determine redox reactions.

The specific objects of current study were to identify the kinetic characteristics and elucidate the possible transformation pathways of redox mediator during the process of microbial goethite reduction. Results from this study give insights into the interaction process of “Quinone-Iron-DIRB” by two specifically designed pre-incubation reaction systems, which were previously unrecognized, in the transformation between oxidized and reduced species of the redox mediator under biotic/abiotic driven

Results

The characteristics of microbial goethite reduction in “strain S12-redox mediator” pre-incubation reaction systems. Both the concentrations of dissolved reductively produced ferrous iron (Fe(II)\text{red}) and total reductively produced ferrous iron (Fe(II)\text{tot}) increased significantly (Fig. 1) in the “strain S12-redox mediator” pre-incubated reactors spiked with goethite of different concentrations. The reductively production of ferrous iron was attributed to the iron reducing bacterial respiration and the abiotic reduction of goethite by the reduced form of the redox mediator (AQS\text{red}). AQS\text{red} was formed during the pre-incubation duration (3 days) (Fig. 1A in Supporting Information (Fig. S1A), from beginning to 3 hours incubation), and it was subsequently transformed to the oxidized form (AQS\text{ox}) after goethite spike (Fig. S1A, from 3 hours to 3.5 hours incubation), indicating the role of AQS in reducing goethite.

In all the reactors spiked with different concentrations of goethite, Fe(II)\text{ox} and Fe(II)\text{dis} concentrations increased with time and then reached a plateau. The incubation time needed to reach the plateau increased with increasing concentration of the spiked goethite. In reactors containing no more than 0.5 mM goethite, the goethite reduction reached the plateau after about 5 days of goethite reduction (Fig. 1A,B). However, in the reactors containing spiked goethite higher than 2.0 mM, it would take 10 days or more incubation time to reach the plateau (Fig. 1C–E).

In “Strain S12- AQS” pre-incubation reactors, a positive linear relationship exists between the mass-normalized goethite reduction rate and goethite mass-available AQS content (Adj. R-Square = 0.9967) (Fig. S2), which could be due to a change in available surface area per unit goethite mass for redox mediator (AQS) during the subsequent interaction process between redox mediator and iron oxide. AQS content was kept constant at 0.1 mM in each “strain S12- AQS” pre-incubation reactor, while goethite concentration varied from 0.1 to 10.0 mM, which changed the value of goethite mass-available AQS content ([AQS]/[Goethite]). In the reactors containing lower goethite content, more AQS is available for goethite surface site, consequently enhancing the overall potential of electron transfer from the reduced form of AQS to goethite, and increasing the mass-normalized goethite reduction rate. This explained why the reaction time needed for Fe(II) concentration could reach a plateau (dotted line in Fig. 1) increased with increasing content of goethite.

Furthermore, the value of [AQS]/[Goethite] also significantly impacted the relative extent of goethite reduction. Almost 100 percent of goethite was reduced in the reactor containing 0.1 mM goethite ([AQS]/[Goethite] = 1) while the value in the reactor containing 10.0 mM goethite ([AQS]/[Goethite] = 0.1) reached 32.1 percent at the plateau (Fig. S3, solid symbol).

Influence of contents of AQS addition to the “strain S12-goethite” pre-incubation reaction systems on microbial goethite reduction. Spiking AQS in the pre-incubation reactors of “strain S12-goethite” resulted in a significant increase in Fe(II)\text{ox} and Fe(II)\text{dis} concentrations, indicating the increase of goethite reduction rate compared with that before AQS addition (gray area in Fig. 2). The goethite reduction rate increased with the increasing concentration of spiked AQS. For example, the average rate of goethite reduction increased by 3 times after spiking 0.3 mM AQS (Fig. S4). Increasing spiked AQS concentration also increased the extent of goethite reduction, and in the reactor containing a relatively high content of AQS (0.30 mM), nearly 85 percent of goethite was reduced; when the AQS concentration decreased to 0.025 mM, only 53 percentage of goethite were reduced (Fig. S3, open symbol). The rate of goethite reduction decreased sharply after nearly 15th days’ incubation (or 8 days after AQS spike regardless of spiked AQS concentration (dotted lines in Fig. 2A–D), which indicated that the microbial reduction of goethite mediated by AQS reached the equilibrium.

The introduction of redox mediator, AQS, could promote the rate (Fig. S4) and extent of the reductive dissociation of goethite (Fig. S3). Thus the following could be expected to occur: a) Fe(II)\text{ox} concentration was increased; b) the enhanced microbially mediated reductive dissolution could result in a smaller size of goethite particle, and subsequently, the specific surface area and adsorption capacity of goethite were enhanced. For example, in “strain S12-goethite” pre-incubation reaction systems containing 0.3 mM AQS, the adsorption capacity of goethite for ferrous iron increased to 1.72 mM·mM\textsuperscript{-1} (Fig. S5), thus, the contents of adsorbed ferrous iron (Fe(II)\text{ads}) being
Figure 1. Concentrations of Fe(II)\textsubscript{dis} and Fe(II)\textsubscript{tot} in “strain S12- AQS” pre-incubation reaction systems added with different content of goethite after 3 days incubation. (Solid symbol: Fe(II)\textsubscript{tot}, open symbol: Fe(II)\textsubscript{dis}). Reaction systems components: AQS: 0.1 mM; goethite: 0.1 ~ 10.0 mM. The reaction systems containing 0.1 mM AQS inoculated with strain S12 at density of 2.5 × 10^8 cells·ml\(^{-1}\), then after 3 days pre-incubation, different contents (0.1, 0.5, 2.0, 5.0 and 10.0 mM) of goethite were added to each pre-incubation reaction systems (A–E) respectively.)
elevated (Fig. 3). The similar phenomenon could also be seen in “strain S12-AQS” pre-incubation reaction systems (Fig. S6).

**Characteristics of redox mediator transformation in two different pre-incubation reaction systems.**
The maximum absorbance wavelength of oxidized form of redox mediator (AQS$_{oxi}$) and its hydroquinone reduced form (AQS$_{red}$) were near 329 nm and 380 nm respectively. Considering that the maximum molar absorption coefficients for AQS$_{oxi}$ and AQS$_{red}$ are constants (Table S1), the absorbances at 329 nm and 380 nm will be directly proportional to the concentration of AQS$_{oxi}$ and AQS$_{red}$ respectively.

In “strain S12-AQS” pre-incubation reactors, redox mediator was gradually transformed to AQS$_{red}$ within initial 3 days of pre-incubation duration (Fig. 4A). After goethite spike, AQS$_{red}$ decreased sharply and was almost completely transferred to its oxidized form. For example, after 0.5 days incubation with addition of 5.0 mM goethite, the absorbance of 380 nm at which the AQS$_{red}$ has the maximum absorbance was hardly detected (dotted line in Fig. 4). Further investigation of the UV/vis spectra during the subsequent reduction of goethite indicated that the concentration of AQS$_{red}$ increased gradually from 3.5 to 14 days (Fig. 4B). The similar phenomenon could also be seen in other “strain S12-AQS” pre-incubation reaction system containing 0.5 mM goethite (Fig. S1).

In “strain S12-goethite” reactors, after AQS$_{oxi}$ was spiked at different concentrations, the transformation from AQS$_{oxi}$ to AQS$_{red}$ was confirmed by UV-vis absorption spectroscopy (Fig. S7). The temporal change in maximum absorbance of the AQS$_{oxi}$ at 380 nm could be described by an exponential growth (ExpGro) model (Fig. 5) with the correlation coefficients, Adj. R-Square, were in the range of 0.7768 ~ 0.9762 (Table S2). The fitted ExpGro model was discussed in Text 1 in Supporting Information (Text S1).
Discussion

From the analysis of the characteristics of transformation between two redox species of redox mediator in "strain S12-AQS" pre-incubation reaction systems, the transformation process mentioned above could be described by the Fig. 6.

It indicated an obvious reversible redox transformation between two species of redox mediator (Fig. 6), AQS\textsubscript{oxi} and AQS\textsubscript{red}, with the detailed processes proposed here: 1) microbial mediation pathway (Reaction І in Fig. 6): the AQS\textsubscript{oxi} received the enzymatically produced electrons and was reduced to AQS\textsubscript{red}, which could be accumulated till the equilibrium of the transformation of two redox sensitive species (from beginning to 3 days, Fig. 4A); 2) abiotic pathway (Reaction ІІ in Fig. 6): when goethite was introduced to the reaction system, the accumulated AQS\textsubscript{red} transferred the electrons to goethite surface so as to facilitate the goethite reduction, and meanwhile, AQS\textsubscript{red} was oxidized to AQS\textsubscript{oxi} (from 3 days to 3.5 days, Fig. 4A). 3) The produced AQS\textsubscript{oxi} was subsequently bio-reduced to AQS\textsubscript{red} during the metabolic process of strain S12 (Reaction ІІІ in Fig. 6) (from 3.5 days to 14 days, Fig. 4B). Therefore, both biotic and abiotic steps controlled the transformation between two species of the redox mediator.
AQS, in the “strain S12- AQS” pre-incubation reaction systems; and consequently, the concentrations of AQS_{red} and AQSoxi varied in different incubation times (Fig. 4).

Further investigation showed that during the transformation process from AQS_{red} to AQSoxi (Reaction II in Fig. 6), the relationship between the initial AQS_{red} oxidization rate and total concentration of added goethite could be fitted by the exponential growth model (Adj. R- Square = 0.9942). From the fitted parameters, the initial AQS_{red} oxidization rate reached a maximum at about 0.22 mM·d^{-1} when goethite concentration was high enough to be at 10.0 mM (Fig. 7).

Figure 5. The times course of the maximum absorbance of reductively produced AQS_{red} at 380 nm in “strain S12- goethite” pre-incubation reaction systems added with different concentration of AQS.

Figure 6. Proposed pathways for the interaction process among AQS, strain S12 and goethite in “strain S12-AQS” pre-incubation reaction systems.

Figure 7. The influence of added goethite contents on the initial AQS_{red} oxidation rate in “strain S12-AQS” pre-incubation reaction systems. (Goethite: 0.1 ~ 10.0 mM; AQS: 0.1 mM; inoculation cell density: 2.5 × 10^8 cells·ml^{-1}).
In “strain S12-AQS” pre-incubation reaction systems with the concentration of redox mediator (AQS) kept constant at 0.1 mM, the value of goethite mass-available redox mediator content ([AQS]/[Goethite]) could be low enough to be at 0.05 or even less in the reaction system containing 2.0 mM goethite or higher. The decreased value of [AQS]/[Goethite] indicated that the redox mediator cannot completely occupy the active surface sites on the goethite surfaces when goethite concentration was increased. Then the influence of goethite concentration on the initial AQS_red oxidation rate became not obvious when goethite concentration was higher than 2.0 mM, [AQS]/[Goethite] < 0.05, which was in contrast to that in reaction systems containing low concentrations of goethite (range from 0.1 ~ 2.0 mM) (gray area in Fig. 7), and thus the typical saturation-type exponential growth (ExpGro) model could be fitted for the relationship between the initial AQS_red oxidation rate and the concentrations of added goethite (Fig. 7).

The similar trend could also be seen in reaction systems with the low inoculation cell density at 5.0 × 10^7 cells·ml⁻¹ (Fig. S8), and however, the initial AQS_red oxidation rate could only reach a maximum at no more than 0.12 mM·d⁻¹ (Fig. S8). Low inoculation cell density was expected to result in decreased microbial activities, and the initial AQS_red oxidation rate decreased consequently. The results of fitting Monod-type equation of initial microbial goethite reduction rates (Fig. S9) further proved that the degree of biotic reaction reached 0.5361 mM·d⁻¹ (inoculation cell density: 2.5 × 10^7 cells·ml⁻¹) which was nearly twice as much as that with low inoculation cell density (5.0 × 10^7 cells·ml⁻¹) (Table S3).

When AQS_red was added to “strain S12-goethite” pre-incubation reaction systems, it received the enzymatically produced electrons and was bio-reduced to AQS_red, which could subsequently transfer the electrons to goethite surface and enhance the goethite reduction (Fig. 2). The interaction process among redox mediator, strain S12 and goethite could be described by the Fig. 8.

The bio-reduced AQS_red was accumulative in the first several incubation days when redox mediator was introduced in “strain S12-goethite” pre-incubation reaction systems (Fig. S7). The result implied that the rate of AQS_red formation by the biotic Reaction III in Fig. 8 would be higher than that of AQS_red consumption by the abiotic Reaction II in Fig. 8, and thus the produced AQS_red through Reaction III exceeded that of AQS_red oxidized to AQS_red through Reaction II, resulting in the accumulation of bio-reduced AQS_red (Fig. 5).

The concentrations of Fe(II)₉ and Fe(II)₈ in “strain S12-goethite” pre-incubation reaction systems increased when different concentrations of redox mediator were introduced, and eventually they leveled off after about 15 days incubation period (Fig. 2), based on Fig. 8, it indicated that the rate of microbial goethite reduction (Reaction I) and redox mediator-assistant abiotic goethite reduction (Reaction II) both decreased and almost got equilibrium. During the course of rate decreasing of Reaction II in Fig. 8, the remaining AQS_oxi in reaction systems would be transformed to AQS_red driven by Reaction III in Fig. 8, and subsequently the content of AQS_red would reach a plateau (Fig. 5) indicating that the transformation between AQS_oxi and AQS_red under biotic driven was in equilibrium.

A spontaneous reaction occurs in the direction of increasing the reduction potential (ΔE). The reduction potentials of Reaction II in Fig. 8, could be calculated using the Nernst equation, allow the energetics of each half-reaction to be evaluated separately (details seen in Text S2):
With increasing content of AQS added to the “strain S12-goethite” pre-incubation reaction systems, the value of \( \frac{[\text{AQS}_{\text{oxi}}]}{[\text{AQS}_{\text{red}}]} \) decreased, while the value of \( \frac{[\text{Fe(II)}]}{[\text{FeOOH}_{\text{goethite}}]} \) increased (Fig. 9). Based on Eq. 1, the value of \( \Delta E \) decreased with the increasing contents of AQS added to reaction system. Since a decreasing reduction potential means that reduction reaction approaches equilibrium and it occurs more difficulty, that is to say, the reduction reaction rate of Reaction II in Fig. 8 decreases consequently with the increasing contents of AQS added to reaction system.

During the course of the reduction reaction rate decreasing of Reaction II in Fig. 8, that is to say, the rate of the AQSred consumed by Reaction II decreased. However, the AQSred was continually produced from Reaction III in Fig. 8, it will accumulate and subsequently the concentration of AQSred would reach a plateau. In each “strain S12-goethite” pre-incubation reaction systems, the inoculation cell density was kept constant and microbial activities could be assumed equal. Thus the concentration of AQS would be the rate limiting factor for biotic Reaction III in Fig. 8 which driven the transformation from AQSoxi to AQSred. Then the average apparent AQSred formation rate through Reaction (III) in Fig. 8 would be expected to increase with increasing concentration of AQS (Fig. 10),
and a positive linear relationship with the correlation coefficients of 0.9937 was found between AQS contents and the average AQS\textsubscript{red} formation rate in “strain S12-goethite” pre-incubation reaction systems.

Organic matters are redox active natural macromolecules that are ubiquitous in soils and sediments and predominate in the subsurface. The redox activity of organic matter has been primarily ascribed to quinone-hydroquinone moieties. In environments with variable redox conditions, organic matters containing a pool of functional moieties have reversible electron transfer ability and can act as redox mediator\textsuperscript{26}. Thus organic matters shuttle electrons from microbe to poorly accessible mineral phases, such as iron oxides, could not only act as redox buffers by accepting electrons from microbial respiration under anoxic condition but also mediate biogeochemical redox reactions\textsuperscript{26}. These results of our current investigation will help to understand the microbial reduction of the Fe(III) oxide, especially in natural environment containing various organic matters, which can act as the potential electron acceptor or donor to facilitate the interaction between microbe and Fe(III) oxides.

**Methods**

The preparation of deoxygenated distilled deionized water, properties of chemicals electrolyte solutions, and anaerobic glove box were provided in Text S3.

**Goethite synthesis and SEM, XRD analysis.** Goethite was used in the experiments as the model Fe(III) oxide since it is commonly found in environment. Goethite was synthesized using a method described by Schwertmann and Cornell\textsuperscript{28}. Details of the syntheses, washing, and characterization of the goethite were provided in the Text S4 and Figs S10 and S11. Briefly, 120 ml of 1 M Fe(NO\textsubscript{3})\textsubscript{3} was mixed with 200 ml of 4.5 M KOH solution, and the mixture was then boiled at 70 °C for 60 hs by constant stirring. After being cooled down to the room temperature, the mixture was filtered, and the yellowish precipitates on the filter were washed repeatedly using deionized water to remove any dissolved salts. The goethite slurries were diluted to 200.0 mM in water and were sonicated for 1 h to disperse the lyophilized particles.

**Redox mediator and its UV-vis spectra.** Anthraquinone-2-sulfonate(AQS) was used as the model redox mediator(Table S1) due to its high water solubility, and the convenience of analysis using UV-visible spectrophotometry. AQS has oxidized and reduced forms as quinone and hydroquinone moieties, respectively. The maximum absorbances of the oxidized form (AQS\textsubscript{ox}) and reduced form (AQS\textsubscript{red}) were at 329 and 380 nm wavelength, respectively. The redox transformation between AQS\textsubscript{red} and AQS\textsubscript{ox} can be determined using UV-vis absorption spectroscopy. Procedures for obtaining absorption spectra of different forms of redox mediator (AQS\textsubscript{red} and AQS\textsubscript{ox}) were reported in our previous study\textsuperscript{6}. Briefly, 1.5 ml suspension in a reactor was transferred to 2 ml centrifuge tube, and centrifuged for 2 minutes at 13000 rpm. The supernatant containing redox mediator was collected by a quartz cuvette and was quickly scanned using a UV-visible spectrophotometer (DR 5000, HACH™, USA).

**Bacteria and medium.** *Shewanella decolorationis* S12 (strain S12), a new species of the genus *Shewanella*, was provided courtesy by Dr. Xu (Guangdong Provincial Key Laboratory of Microbial Culture Collection and Application, Guangzhou, China). Strain S12 was isolated from the activated sludge of a textile printing wastewater treatment plant in Guangzhou, China. Strain S12 can grow in the presence of various electron acceptors, such as oxygen, nitrate, nitrite, ferric iron and sulfate, showing remarkable respiratory versatility\textsuperscript{29}, as do other members of DIRB\textsuperscript{30,31}.

Strain S12 was routinely cultured aerobically at 32 °C in Luria-Bertani (LB) medium. The LB medium contains tryptone(10.0 g·L\textsuperscript{-1}), yeast extract(5.0 g·L\textsuperscript{-1}), NaCl(10.0 g·L\textsuperscript{-1}) at pH 7.0 (adjusted with 5 M NaOH), and was sterilized by autoclaving for 30 mins. The strain was harvested from the LB cultures at mid to late log phase by centrifuging the suspension for 10 mins at 5 °C. The cells were washed with a pH buffer (1.4- piperazinediethanesulfonic acid (PIPES) buffer, 15.0 mM, pH 7.02) to remove residual medium, and resuspended in the buffer.

**Experiment design and procedures.** In natural environment, it is expected that indigenous bacteria and Fe(III) oxide minerals(electron acceptors) or redox sensitive humic substance(redox mediator) are co-present for an extended period of time prior to the interaction among “redox mediator-iron oxide-DIRB”. Based on these considerations, the following sets of experiments were designed and performed:

1. “strain S12- redox mediator” was first incubated in solutions containing strain S12 and 0.1 mM AQS for 3 days, and this incubated system was then spiked with goethite at variable concentrations (0.1 ~ 10.0 mM) to examine the possible transformation pathway of AQS during the interaction process of AQS-goethite.

2. “strain S12-goethite” was first incubated in solutions containing strain S12 and 2.0 mM goethite for 7 days, and the incubated system was then spiked with AQS\textsubscript{ox} at variable concentrations (0.025 ~ 0.30 mM) to evaluate subsequent AQS-goethite interactions.

In a typical experiment, various volumes of goethite slurry and lactate were mixed to achieve desired concentrations, followed by addition of PIPES buffer solution (15.0 mM, pH 7.02), AQS (0.1 mM) and NH\textsubscript{4}Cl (20.0 mM). Tubes were purged with N\textsubscript{2} and sealed with thick butyl rubber stoppers. Strain S12 cell suspension (D600 nm = 1.75) was added from a freshly washed culture in PIPES buffer system using a needle and syringe.

**Definition of different species of reductively produced ferrous iron.** The suspension samples were taken at selected times, being filtered to measure dissolved Fe(II) using the ferrozine method\textsuperscript{32}. The total Fe(II)
was measured by acid-extraction (0.5 M HCl) for one hour. The extracted Fe(II) was then measured by the ferrozine method. The difference between total and dissolved Fe(II) is the sorbed Fe(II) in the suspensions.

For the convenience of discussion, different species of reductively produced ferrous iron were defined as Eq. 2:

$$Fe(II)_{tot} = Fe(II)_{dis} + Fe(II)_{ads}$$  

(2)

where \(Fe(II)_{dis}\) is dissolved Fe(II) as measured in filtered (0.45 μm) samples, \(Fe(II)_{tot}\) is the total Fe(II) extracted by 0.5 M HCl, and \(Fe(II)_{ads}\) is the adsorbed ferrous iron calculated from the difference of the Fe(II)\text{dis} and Fe(II)\text{tot}. All the procedures were performed in an anoxic chamber (Bactron III, SHELLAB™, USA).

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Author Contributions

W.Z. designed experiments and analysed data, M.S. and D.Y. performed experiments. C.L., T.H. and F.W. prepared the manuscript.
Additional Information

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