
Anonymous Referee #2

General Comments:

The manuscript is well written and the main points come across easily. In the introduction, the authors recognize a number of studies that analyze the influence of organic matter (OM) on Fe(III) reduction kinetics and Fe-mineral transformations. These studies have been carried out under abiotic conditions or with pure cultures of well-known Fe-reducers such as Geobacter and Shewanella. This study focuses on (1) defining how a natural microbial consortium influences the reduction of Fe(III)-OM complexes and (2) determining microbial community changes under anaerobic Fe-reducing conditions. These findings will significantly impact our understanding of Fe-OM interactions under environmentally relevant conditions.

We thank our second referee, anonymous referee #2, for taking the time to review our manuscript and for his/her feedback and helpful comments. Your comments will definitely enhance the quality of the manuscript. Below you will find our responses to all comments.

Specific Comments:

Introduction

Line 33: should be “The majority of which is dispersed..”
Corrected (Line 33).

Line 44: should be “The coprecipitation of OM with Fe results in...”
Corrected (Line 44).

Line 49-51: This sentence is confusing. It seems like you are trying to make two separate arguments (1) coprecipitated ferrihydrite has different properties than pure ferrihydrite, and (2) coprecipitated ferrihydrite has different reactivity compared to ferrihydrite with sorbed OC. Maybe split into two sentences?

Sentence revised for clarification as follows: ‘The different properties of pure ferrihydrite and OM-ferrihydrite coprecipitates may lead to different behaviors during microbial reduction. Due to their smaller crystal size and more defective crystal structure, coprecipitates might faster dissolve. The associated organic material will change the mineral’s surface properties, e.g., the surface charge, with consequences for the accessibility of Fe(III) to microbes, redox-active shuttling compounds, or extracellular enzymes.’ (Line 49-52).

Line 74-93: This paragraph has important information that you reply on in your discussion. Lines 94-97 clearly state the overall findings, but I found it a bit difficult to follow lines 74-93.

Modifications to this section were included to improve the flow (Line 94-101)
Line 101: should be “is the dominant electron accepting process”
Corrected (Line 104).

Methods

Line 110-112: This first sentence is confusing.

Sentence modified for clarification as follows: ‘Peat cores were obtained from the Schlöppnerbrunnen fen (Northern Bavaria, Germany; 50°07′55″N, 11°52′52″E) using a Pürkheimer soil corer in May 2016. This minerotrophic, slightly acidic (pH ~5) fen has been previously described in detail (Blodau et al., 2004; Eusterhues et al., 2014; Hausmann et al., 2016; Küsel et al., 2008; Loy et al., 2004; Pester et al., 2012).’ (Line 119-121).

Line 117: what is Corg?

‘Solid organic carbon’ added for clarification (Line 127).

Line 129: This first sentence seems like it should be last sentence of the previous paragraph.
The sentence is now the concluding sentence of the previous paragraph as suggested (Line 139-140). Line 130-133: These sentences are out of place. I don’t think you need to talk about DNA extraction or PCR here. Maybe move them to the “DNA extractions” and “Quantitative PCR” sections.

The descriptions regarding DNA extraction and subsequent qPCR analyses were moved to section 2.5 (Line 196-202).

Line 146: Move this up so that the reader knows that you performed the forest floor extract solutions. Information about forest floor extract solution moved to Line 144 (detailed descriptions can be found in Esterhues, et al. 2014).

Line 149-156: You may want to add this paragraph to a section called “Preparation of pure cultures”. We appreciate the suggestion, however, we have decided to keep the information regarding pure culture preparation and the subsequent incubation set-up with both S. oneidensis and the microbial consortia extracted from peat in one section instead of separating the information into separate sections (2.4 Microbial OM ferrihydrite reduction experiments).

Sentence modified to prevent repetition of detailed description as follows: ‘Cells were washed twice with a defined medium, resuspended in 2 ml of medium and diluted to a final concentration of 2 x 10^5 cells ml^-1 in culture tubes, containing a defined medium.’ (Line 168-170).

Line 176: If you are not showing any data (not even in the supplemental information), then why mention that you did this?

Secondary mineral formation during reduction is of interest in soil and sediment research. Many previous studies address this question. We therefore believe it is important to mention that we also paid attention to this aspect, although we did not find any new minerals. However, a table or a figure with 21 very similar diffractogramms does not seem necessary to us to document our findings.

179: Did you also collect XRD spectra of the starting materials?

Yes, XRD spectra was collected from the starting material. Section 2.10 (Line 264-265) and 3.4 (Line 360-361) were both modified to reflect this.

Line 199: move this sentence to the end so that it follows the order in which things were done. It will also make it easier for the reader to understand that you did the PCR and prepared the libraries yourself and just did the sequencing at the LGC Genomic GmbH.

PCR, library preparation, and sequencing was all done by LGC Genomic GmbH. Including this type of detailed information (i.e. primers, etc.) in the materials and methods is commonly done. Sentence added to this section to make it clear that LCG did the PCR, library prep, and subsequent sequencing.

Line 224: Did you normalize the data using qiime or any other method? If you did, you should mention this because it is important when comparing the abundance of different taxa across various samples.

Yes, the data was normalized using qiime. An additional sentence was added to explain how differentially abundant taxonomic groups were identified after the metadata was assembled for downstream analysis in qiime (Line 252-256).

Results The order of these sections has a great flow. However, I would consider merging the Results and Discussion sections into one section. This will avoid many repetitive statements.

We opted to keep the results and discussion sections as independent section and based on comments you provided as well as the other reviewer we hope the repetitiveness is minimized while maintaining the flow of the results and subsequent discussion section.

Section 3.1: The reduction rates stated in this paragraph are not the same as the ones in Table 1.

Reduction rates stated in paragraph corrected to reflect those found in Table 1.

Section 3.3: When reading this section and flipping to the figures, I found it difficult to determine if I was looking at the correct figure.

Please double check the Figure numbers that you refer to in the text. For example, in line 285 you mention beta diversity and Figure 5b and I think you meant to reference Figure 6b.
Figure numbers double-checked. Thanks for pointing this out. Additionally, it’s important to note that all figures were re-numbered due to merging of Figures 1 and 2, 3 and 4.

Discussion
Line 373: Add reference.

References added (Line 452).
Line 394: ‘:’ should be ‘.’

Corrected (Line 473).

References added (Line 486).
Line 435: Have you considered removing singletons and see what the data looks like?

Yes, we did consider this and added now the following sentences to section 2.8 (Microbiome profiling): ‘After alpha diversity analyses, the dataset was stripped by removing OTUs that were represented by less than three sequences. Relative abundances were calculated and standardized based on median sequencing depth across all samples. Datasets were not rarefied to prevent a loss of sensitivity in downstream sample-wise comparisons (McMurdie and Holmes, 2014). The dataset was stripped by removing OTUs that were only identified in single samples (i.e. singletons).’ (Line 243-247). Additionally, the impact on the inclusion of singletons is addressed in Lines 513-515.

Section 4.3: Include mineral formulas and consider citing the following paper: Influence of Coprecipitated Organic Matter on Fe2+(aq)-Catalyzed Transformation of Ferrihydrite: Implications for Carbon Dynamics Chunmei Chen, Ravi Kukkadapu, and Donald L. Sparks Environmental Science & Technology 2015 49 (18), 10927-10936 DOI: 10.1021/acs.est.5b02448

Mineral formulas added to the list of minerals observed in XRD spectra (Line 360-363). Citation also added as suggested.
Line 480: Where is the evidence of this finding? Maybe consider placing this information in the section about mineral composition.

We introduced this information Section 4.1 (Line 447-453) and references were added to support this claim.

Line 500-503: This last sentence is not clear.

The last sentence was rewritten for clarity as follows: ‘This study also revealed that natural soil OM on the surface of ferrihydrites provides enough redox active groups to maintain microbial Fe(III) reduction processes, instead of passivating the surface. While at the same time the OM enhanced overall microbial growth and shaped the microbial community structure.’ (Line 580-585).

References
Consider citing more recent studies such as: Influence of Coprecipitated Organic Matter on Fe2+(aq)-Catalyzed Transformation of Ferrihydrite: Implications for Carbon Dynamics Chunmei Chen, Ravi Kukkadapu, and Donald L. Sparks Environmental Science & Technology 2015 49 (18), 10927-10936 DOI: 10.1021/acs.est.5b02448


Weinan Pan, Jinjun Kan, Shreeram Inamdar, Chunmei Chen, Donald Sparks, Dissimilatory microbial iron reduction release DOC (dissolved organic carbon) from carbon-ferrihydrite association, Soil Biology and Biochemistry, Volume 103, 2016, Pages 232-240, ISSN 0038-0717,
The first two suggested references (Chen, et al. and Adhikari, et al.) were added to the manuscript. Thank you for pointing out that they were missing.

Figures: Consider merging Figures 1 and 2 into a 4-panel figure. This will allow the reader to compare the results obtained from pure cultures and natural inoculum.

Figures 1 and 2 combined into a single 4-panel figure (see Figure 1). Additionally, we modified the figure caption to reflect the information presented in the merged figure.

Consider merging Figures 3 and 4 into a 4-panel Figure (make sure the y-axis is the same for all). This will allow the reader to compare the results obtained from pure cultures and natural inoculum. You may also want to consider changing T0, T middle and Tend to day 0, day X and day 288. This will keep the axis consistent with the other figures.

Figures 3 and 4 combined into a single 4-panel figure (see Figure 2). Figure caption modified to reflect information presented in the merged figure.

Figure 5C. The results illustrated in this Figure were not clear to me. Perhaps state the taxonomic level of the x-axis? Or add different shapes to differentiate between CFh vs Fh and AFh vs Fh?

The taxonomic level is for each portion of Figure 3 (previously Figure 5) is described in the figure caption (i.e. 3A – prokaryotic phyla level; 3B – family level; 3C – genus level).

Figure 6. I understand the importance of using various alpha diversity parameters. However, I am not sure illustrating them all in a main figure is necessary. Perhaps choose one parameters and talk about that one in the paper. The rest could go in the supplemental materials.

This figure was reconfigured to only include Observed OTUs, Chao1 and Shannon indices for simplicity (Fig. 4A). The main text (Section 3.3) and figure caption was modified to reflect these changes.