



Cite this: *Environ. Sci.: Processes Impacts*, 2023, 25, 1945

Contact with soil impacts ferrihydrite and lepidocrocite transformations during redox cycling in a paddy soil†

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Iron (Fe) oxyhydroxides can be reductively dissolved or transformed under Fe reducing conditions, affecting mineral crystallinity and the sorption capacity for other elements. However, the pathways and rates at which these processes occur under natural soil conditions are still poorly understood. Here, we studied Fe oxyhydroxide transformations during reduction–oxidation cycles by incubating mesh bags containing ferrihydrite or lepidocrocite in paddy soil mesocosms for up to 12 weeks. To investigate the influence of close contact with the soil matrix, mesh bags were either filled with pure Fe minerals or with soil mixed with ⁵⁷Fe-labeled Fe minerals. Three cycles of flooding (3 weeks) and drainage (1 week) were applied to induce soil redox cycles. The Fe mineral composition was analyzed with Fe K-edge X-ray absorption fine structure spectroscopy, X-ray diffraction analysis and/or ⁵⁷Fe Mössbauer spectroscopy. Ferrihydrite and lepidocrocite in mesh bags without soil transformed to magnetite and/or goethite, likely catalyzed by Fe(II) released to the pore water by microbial Fe reduction in the surrounding soil. In contrast, ⁵⁷Fe-ferrihydrite in mineral-soil mixes transformed to a highly disordered mixed-valence Fe(II)–Fe(III) phase, suggesting hindered transformation to crystalline Fe minerals. The ⁵⁷Fe-lepidocrocite transformed to goethite and small amounts of the highly disordered Fe phase. The extent of reductive dissolution of minerals in ⁵⁷Fe-mineral-soil mixes during anoxic periods increased with every redox cycle, while ferrihydrite and lepidocrocite precipitated during oxic periods. The results demonstrate that the soil matrix strongly impacts Fe oxyhydroxide transformations when minerals are in close spatial association or direct contact with other soil components. This can lead to highly disordered and reactive Fe phases from ferrihydrite rather than crystalline mineral products and promoted goethite formation from lepidocrocite.

Received 21st July 2023
Accepted 16th October 2023

DOI: 10.1039/d3em00314k

rsc.li/espi

Environmental significance

Iron mineral dynamics are closely coupled to nutrient and contaminant cycling in intermittently or permanently flooded soils, where sub- or anoxic conditions establish frequently. In turn, interactions of Fe minerals with these elements can influence the mineral stability. We show that a close spatial association with the soil matrix drastically impacts Fe mineral transformations. This leads to less crystalline transformation products and increased reductive dissolution with repeated redox cycles. The findings contribute to an improved understanding of Fe mineral dynamics in redox-active soils, such as paddy soils and are important to predicting the biogeochemical cycling of nutrients.

Introduction

When oxygen is depleted under flooded soil conditions, ferric iron (Fe(III)) can act as an alternative electron acceptor during

microbial respiration.^{1,2} This leads to the reductive dissolution of Fe(III) oxyhydroxides,¹ such as lepidocrocite and ferrihydrite, and the release of ferrous iron (Fe(II)) to the soil solution. Since Fe oxyhydroxides are important sorbents for, for example, organic matter, nutrients and trace elements, these compounds can be released into the soil porewater during reductive dissolution.^{3–5} When the soil water content decreases due to drainage or evapotranspiration, the soil becomes aerated and dissolved and/or sorbed Fe(II) is oxidized. This results in the precipitation of mixed-valence or Fe(III) oxyhydroxides and the immobilization of components which coprecipitate or sorb to these phases.^{4,6,7} Therefore, Fe mineral dynamics can control the cycling of, for example, phosphate,^{5,8} arsenic,³ or zinc⁴ in

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d3em00314k>



soils and sediments that are intermittently flooded, such as rice paddy soils.^{9,10}

Under Fe-reducing conditions, Fe(III) oxyhydroxides (hereafter Fe oxyhydroxides) and Fe(II) can coexist. Sorption of Fe(II) to Fe oxyhydroxide surfaces leads to the oxidation of Fe(II) and, through electron transfer, the reduction of structural Fe(III), which is subsequently released as dissolved Fe(II).¹¹ Since ferrihydrite and lepidocrocite are thermodynamically metastable, this process can accelerate their transformations to more crystalline Fe minerals, such as goethite or magnetite.¹² In mineral suspensions, interactions of minerals with Fe(II) result in abiotic ferrihydrite transformation to lepidocrocite, goethite, or magnetite.¹² Lepidocrocite often transforms to magnetite^{13–15} or goethite,^{12,16} or undergoes recrystallization¹⁴ during the interaction with Fe(II). The trajectory of abiotic Fe(II)-catalyzed ferrihydrite and lepidocrocite transformation depends on the Fe(II)–Fe(III) ratio^{12,14,17} and the pH of the solution.^{15,18,19} For example, magnetite formation from Fe(II)-catalyzed ferrihydrite transformation is favored at high Fe(II)–Fe(III) ratios and pH > 7,^{12,14} while slightly acidic pH conditions (6–7) favor lepidocrocite and goethite formation.^{12,14} When transformations of ferrihydrite and lepidocrocite are mediated by dissimilatory Fe-reducing bacteria, the biotic mineral transformations can lead to the formation of ferrous carbonate minerals, such as siderite (FeCO₃),²⁰ and green rust²¹ in addition to magnetite and goethite.^{20–22} Abiotic and biotic transformations of Fe oxyhydroxides can be hindered or altered by the presence of, for example, organic matter,^{23–25} trace metals,²⁶ or other oxyanions.^{14,27–29} For example, the presence of organic matter^{23,24} and silicate^{14,27} can hinder the abiotic Fe(II)-catalyzed transformation of ferrihydrite to goethite and magnetite.

Model studies, like those mentioned above, used mineral suspensions or flow-through columns to demonstrate the effect of individual components on Fe mineral transformations. Yet, Fe oxyhydroxide transformations in natural systems are influenced by soil porewater, which comprises complex mixtures of dissolved organic matter, nutrients and trace elements. Previous works aimed at studying mineral transformations in soil and incubated ferrihydrite in flooded soils using permeable bags^{30,31} or diffusive gel samplers.³² In contrast to these studies, where ferrihydrite was spatially separated from the soil matrix and dissolved soil components, including Fe(II), were required to diffuse into the permeable bags or sample holders,^{30–32} Fe minerals in natural systems are part of the soil matrix and thus, are more closely associated to or in direct contact with organic matter, other minerals, and microorganisms. This likely impacts the trajectory of mineral transformations.³³

Redox conditions in intermittently flooded soils may alternate between oxic and sub- or anoxic conditions. Therefore, several studies have investigated Fe mineral dynamics in soils during redox cycles,^{34,35} mainly using soil slurries³⁶ purged with N_{2(g)} or O_{2(g)}.^{37,38} After exposure to one or multiple redox cycles, both an overall increase^{34,35,37} and decrease^{35,37,38} in short-range-ordered Fe minerals in the soil has been reported. Changes in the short-range-ordered Fe mineral

fraction in soils can strongly impact the bioavailability of Fe for microbial reduction and thus the Fe(II) production upon the onset of Fe reducing conditions.³⁹ Short-range-ordered Fe phases, such as ferrihydrite, are regularly precipitated during oxic periods of redox cycles^{36–38} and are highly susceptible to mineral transformation.^{12,21} During redox cycles, reductive Fe mineral transformations are interrupted by oxic periods, which may lead to different mineral transformation extents compared to mineral transformations under permanently sub- or anoxic conditions.

Due to the importance of Fe minerals for element cycling, we need to improve our understanding of how Fe oxyhydroxides transform in soils. In comparison with mixed soil slurries used in previous studies,^{36–38} undisturbed soils are characterized by higher soil-to-solution ratios, chemical and physical heterogeneity at the pore- and aggregate scale,^{40–44} and pronounced diffusion limitations of chemical processes.^{40,41} These factors likely influence Fe mineral transformations, for example, by impacting the local concentrations of Fe(II) and organic matter. To better understand Fe mineral dynamics in soils, recent approaches combined the use of ⁵⁷Fe-labeled solutions or minerals in soil experiments with ⁵⁷Fe Mössbauer spectroscopy, which is only sensitive to ⁵⁷Fe.^{33,45,46} For example, Tishchenko *et al.*⁴⁶ reacted ⁵⁷Fe(II) with a soil slurry and used Mössbauer spectroscopy to study the Fe atom exchange between dissolved Fe(II) and structural Fe(III) and Fe mineral crystallization. Chen and Thompson⁴⁵ added dissolved ⁵⁷Fe(II) to soils, oxidized the soil slurries, and analyzed the oxidation products with Mössbauer spectroscopy. Recently, Notini *et al.*³³ demonstrated a new approach for studying the transformations of synthetic Fe minerals in close contact with soil by using ⁵⁷Fe-labeled minerals and following their transformation with Mössbauer spectroscopy. Adopting this approach in the current study, we follow the transformation of ⁵⁷Fe-labeled ferrihydrite and lepidocrocite in a soil matrix exposed to redox fluctuations.

The objectives of this study were to (i) investigate how redox fluctuations affect the transformations of ferrihydrite and lepidocrocite in a paddy soil at a natural soil-to-solution ratio and (ii) to assess how the close spatial association between the added Fe minerals and the soil matrix affects the Fe mineral transformations. We incubated ferrihydrite and lepidocrocite in soil mesocosms exposed to three flooding–drainage cycles. As a control, an additional mesocosm remained permanently flooded. Ferrihydrite and lepidocrocite were incubated in mesh bags either containing the minerals without soil or ⁵⁷Fe-labeled minerals mixed with soil to create a close spatial association with the soil matrix. The mineral transformation products in the incubated samples were analyzed with Fe K-edge X-ray absorption fine structure spectroscopy and/or X-ray diffraction and ⁵⁷Fe Mössbauer spectroscopy.

Materials and methods

Soil sampling and characterization

A rice paddy soil was sampled at the Ubon Ratchathani Rice Research Center in Thailand during the dry season in February



2020. A soil profile of 2 m depth was established in the dry rice field and classified as a Hydragric Loamic Anthrosol on sandstone (details in the ESI, Section S1†) after the World Reference Base for Soil Resources.⁴⁷ Surface soil (0–15 cm depth, 100 kg) was sampled for the incubation experiments and was oven-dried at 30 °C until constant weight and homogenized by mixing and sieving (<2 mm). The texture of the sieved soil was silty sand (3% clay, 12% silt, 85% sand). For chemical and mineralogical analyses, representative subsamples of the sieved soil were milled with a vibratory disc mill. The total element contents were measured with X-ray fluorescence spectrometry (XRF; XEPOS, Spectro, Germany) in the milled soil (Fe = 3.3 g kg⁻¹, aluminum = 14.0 g kg⁻¹, silicon = 417.4 g kg⁻¹, Table S1†). Carbon (4.0 g kg⁻¹) and nitrogen (0.5 g kg⁻¹) contents were determined using a CN elemental analyzer (vario MAX cube, Elementar, Germany). The pH of the soil was weakly acidic (pH 5.5 in 0.01 M CaCl₂).

Mineral synthesis

Ferrihydrite and lepidocrocite with natural abundance (NA) Fe isotope composition (⁵⁴Fe-Fh, ⁵⁴Fe-Lp) were synthesized by adapting the methods of Schwertmann and Cornell.⁴⁸ Briefly, for ⁵⁴Fe-Fh synthesis, a 0.2 M ferric nitrate (Fe(NO₃)₃·9H₂O, Merck) solution was titrated (836 Titrand, Metrohm) with 1 M NaOH (Titrisol®, Merck) to pH 7.5 ± 0.1 under vigorous stirring at room temperature. For ⁵⁴Fe-Lp, a 0.2 M Fe(II) solution prepared from FeCl₂ (Merck) was titrated to pH 6.7–6.9 with 1 M NaOH at room temperature, then oxidized under vigorous stirring and gentle purging with air (approx. 100 mL min⁻¹) while the pH was maintained by the further addition of 1 M NaOH. The isotopically labeled ferrihydrite (⁵⁷Fe-Fh) and lepidocrocite (⁵⁷Fe-Lp) were prepared by dissolving ⁵⁷Fe(0) (96.14% ⁵⁷Fe, Isoflex USA) in 2 M HCl (NORMATOM®, 34–37%, VWR) overnight, resulting in a 0.18 M ⁵⁷Fe(II) solution. For ferrihydrite synthesis, the ⁵⁷Fe(II) solution was oxidized to ⁵⁷Fe(III) by adding 35% H₂O₂ (Merck). The ⁵⁷Fe(II) and ⁵⁷Fe(III) solutions were passed through a 0.45 μm nylon filter before ⁵⁷Fe-Fh and ⁵⁷Fe-Lp were prepared as described for ⁵⁴Fe-Fh and ⁵⁴Fe-Lp. All precipitates were repeatedly washed by resuspending the precipitates in ultra-pure water (UPW, ≥18.2 MΩ·cm, Milli-Q, Millipore), centrifuging the suspensions (3438 g for 15 min, 20 °C) and decanting the supernatant. After the washing procedure, the precipitates were again resuspended in UPW, shock frozen dropwise in liquid N₂, freeze-dried, gently homogenized with mortar and pestle, and stored in amber glass bottles in a desiccator. All minerals were characterized by X-ray diffraction (XRD, Bruker D8 Advance), which confirmed the expected mineral composition and showed no crystalline mineral impurities. However, Mössbauer spectra of the ⁵⁴Fe- and ⁵⁷Fe-lepidocrocites collected at 77 K and 5 K indicated the presence of minor amounts (7% in ⁵⁴Fe-Lp and 16% in ⁵⁷Fe-Lp) of nano-goethite (ESI Section S2†). Mineral characterization details, including XRD patterns (Fig. S3†) and Mössbauer spectra (Fig. S4†) with fit components (Table S2†) are provided in ESI Section S2.†

Preparation of mesh bags and sample holders

For the incubation of the minerals, mesh bags (internal dimensions ~1 × 3 × 0.2 cm) were prepared from a polyethylene terephthalate (PETE) mesh fabric (SEFAR, Switzerland). The mesh fabric had a pore size of 51 μm to allow the free exchange of water, solutes and bacteria. The mesh fabric sheets were triple-layered, folded, and heat-sealed on two sides before filling. For the ⁵⁴Fe-minerals, 100 mg of ⁵⁴Fe-Fh or ⁵⁴Fe-Lp were filled inside each mesh bag before the third side was carefully heat-sealed. For the ⁵⁷Fe-minerals, 8 mg of ⁵⁷Fe-Fh or ⁵⁷Fe-Lp were thoroughly mixed with 800 mg of the sieved soil in a 2 mL Eppendorf tube using a plastic rod before being transferred to the mesh bags, as described above. The mineral-to-soil ratio in the mineral-soil mixes was chosen to achieve minimal addition of Fe to the soil while ensuring a sufficiently high Mössbauer signal contribution from the ⁵⁷Fe-labeled minerals, as discussed in Notini *et al.*³³ In this experiment, the addition of the Fe minerals tripled the total soil Fe content while >98% of ⁵⁷Fe in the mineral-soil mixes originated from the added ⁵⁷Fe-labeled minerals. Sample holders were designed and 3D-printed (Clear Photopolymer Resin, Formlabs) to have a pointy tip to enable the insertion of mesh bags into the soil. The sample holders had large vertical openings on the side to maximize the contact between the mesh bag and the surrounding soil. The mesh bags were placed in the sample holders and a threaded plastic rod was attached to facilitate the insertion into the soil (ESI Section S3†).

Mesocosm setup and flooding of the soil

The soil mesocosms were designed to mimic flooding and drainage cycles in rice paddy soils. Each mesocosm was made from a rectangular plastic box (polypropylene, 37.6 × 26 × 28.3 cm, 20 L). The box had holes (diameter 0.5 cm, *n* = 5) in the bottom to allow the drainage of the mesocosms. A layer of mesh fabric, similar to the mesh fabric used for the mesh bags but with a larger pore size (105 μm), was placed on the bottom of the box to prevent loss of soil during the drainage. The mesh fabric was then covered by a ~1 cm thick sand layer (grain size 0.1–0.5 mm, acid-washed, 1.5 kg sand), followed by 12 kg of sieved soil, resulting in a soil depth of 12 cm. Each mesocosm was contained within a second rectangular plastic box equipped with a drainage port to control the water level in the soil. More details on the mesocosm setup are provided in ESI Section S3.†

The experiment was conducted in a climate chamber at 30 °C air temperature and 50% relative humidity. Each mesocosm was flooded with 8 L of a solution prepared from CaCl₂ and NaCl (1 mM Ca, 4 mM Na, 6 mM Cl) using a peristaltic pump and Tygon tubes (Saint-Gobain Tygon R3607, inner diameter 1.85 mm), with four tubes per mesocosm. The solution composition was chosen based on analyses of porewaters from a similar paddy field at the Ubon Ratchathani Rice Research Center (1.2 mM Ca, 6.8 mM Na). The mesocosms were flooded dropwise (~7 mL min⁻¹ per mesocosm) over 20 hours, during which the position of the tubes was changed multiple times to enable homogenous wetting of the soil and to minimize the



formation of air pockets. Four identical mesocosms were prepared, of which three were designated to undergo water table fluctuations (three weeks flooded, one week drained) to induce soil redox cycles. The fourth mesocosm remained permanently flooded for the duration of the experiment (12 weeks) to compare mineral transformation processes under permanently sub- or anoxic soil conditions to transformations during redox cycles. Flooding-drainage cycles will be referred to as redox cycles and flooded and drained periods will be referred to as anoxic and oxic periods, respectively.

After the first flooding, each mesocosm with fluctuating water table was equipped with one gel-filled redox glass electrode (Mettler Toledo; 3 M KCl, Ag/AgCl reference electrode), one tensiometer (METER group), and three Rhizon porewater samplers (Rhizosphere Research Products, The Netherlands, 0.6 μm pore cut-off) installed at ~ 8 cm depth. The oxidation-reduction potential was recorded hourly using a mobile logger system (pH Meter 913, Metrohm, built-in data logger) and converted (+209 mV, 30 $^{\circ}\text{C}$) to redox potentials relative to the standard hydrogen electrode (Eh). The soil matric potential was recorded every 10 min (ZL6, METER Group). The sample holders were installed in the soil at 8 cm depth. In each mesocosm with fluctuating water table, six sample holders of each mineral ($^{57}\text{Fe-Fh}$, $^{57}\text{Fe-Fh}$, $^{57}\text{Fe-Lp}$, $^{57}\text{Fe-Lp}$) were installed, resulting in 24 samples per mesocosm. In the permanently flooded mesocosm, three sample holders of each mineral ($^{57}\text{Fe-Fh}$, $^{57}\text{Fe-Fh}$, $^{57}\text{Fe-Lp}$, $^{57}\text{Fe-Lp}$) were installed (total: 12 samples). A schematic setup of the experiment is provided in Fig. S5.†

Anoxic sampling and drainage

After soil flooding, the porewater of all mesocosms was sampled at 1, 3, 7, 14 and 21 days, using the three permanently installed Rhizons in each mesocosm. Each Rhizon was connected to a 10 mL pre-acidified plastic syringe to stabilize Fe species immediately. Approximately 10 mL of porewater was sampled and subsequently passed through a 0.45 μm nylon filter for DOC analysis (DIMATOC 2000, Dimatec) or a 0.22 μm nylon filter for total element analysis by inductively coupled plasma optical emission spectroscopy (ICP-OES, Agilent 5100). Porewater element concentrations will be shown as the average of the triplicate porewater samples for each mesocosm in figures and reported as the average of the three redox fluctuating mesocosms (*i.e.* 9 porewater samples) in the text. The pH in the mesocosms was measured manually in the wet soil at sample depth (8 cm) at each porewater sampling with a pH glass electrode (Metrohm).

At 21 days after flooding, one set of mineral samples ($^{57}\text{Fe-Fh}$, $^{57}\text{Fe-Fh}$, $^{57}\text{Fe-Lp}$, $^{57}\text{Fe-Lp}$) was removed from each redox fluctuating mesocosm. The threaded plastic rod was unscrewed, and the sample holder was placed into a 50 mL plastic centrifuge tube. The tube was immediately flushed with $\text{N}_2(\text{g})$ and closed to prevent sample oxidation. All samples were transferred to an anoxic glovebox (N_2 atmosphere; MBraun, <5 ppm O_2) after sampling. In the glovebox, the mesh bags were removed from the sample holders and air-dried for two days

before the samples were homogenized with a mortar and pestle. After the mineral sampling, all redox fluctuating mesocosms were drained by opening the valve of the drainage tube at the bottom of the mesocosms. The drainage of the soil to field capacity (*i.e.* matric potential = -6 kPa) was complete within three days, while residual water remained in the lower boxes of the mesocosms (~ 0.9 L).

Oxic sampling and re-flooding

Seven days after drainage, another set of mineral samples was extracted from each redox fluctuating mesocosm, representing the end of one complete redox cycle. The mesh bags were removed from the sample holders, and the samples were air-dried in the climate chamber for two days before the samples were homogenized using a mortar and pestle. To make sure that air drying did not affect the Fe speciation in the samples, we ran a test incubation where $^{57}\text{Fe-Fh}$ -soil mixes were incubated in the experimental soil for one redox cycle (3 weeks flooded, 1 week drained) and dried under ambient air or in the glovebox. The test samples were analyzed with Mössbauer spectroscopy and showed similar solid-associated Fe(II) fractions (Fig. S7†). After the mineral sampling, all redox fluctuating mesocosms were re-flooded as described above. Due to residual water in the soil and in the lower plastic box, the re-flooding only required 4.8 L of solution per mesocosm to achieve the same water level as in the previous flooded period. The entire procedure, including the anoxic and oxic sampling described above, was repeated two additional times, resulting in redox cycles I–III over 12 weeks.

Anoxic sampling in the permanently flooded mesocosm

The porewater in the permanently flooded mesocosm was sampled on the same days as in the redox fluctuating mesocosms. The pH was measured as described above and the oxidation-reduction potential was measured manually with a redox electrode (Mettler Toledo) at each porewater sampling. The mesh bags in the permanently flooded mesocosm were sampled only at the end of the experiment (12 weeks) following the same procedure as described above for anoxic mineral samplings in the redox fluctuating mesocosms. In the permanently flooded mesocosm, soil surrounding one of the three replicates of $^{57}\text{Fe-Fh}$ and $^{57}\text{Fe-Lp}$ was additionally sampled with a plastic Humax core sleeve (diameter 5 cm) by taking a soil core containing the sample holders. All samples and the soil cores were immediately flushed with N_2 and transferred to the glovebox after sampling. The soil cores were set up vertically in the glovebox, and the water was allowed to drain for one hour before the plastic sleeve was opened. The cores were sampled circularly around the sample holder (three layers of ~ 0.5 cm) using a spatula, and samples were dried in the glovebox. After sampling the soil in the cores, the mesh bags were removed from the sample holders, and the samples were dried and homogenized in the glovebox as described above. Aqua regia digestions of soil core samples combined with Fe isotope analysis of aqua regia extracts (methods described below) allowed us to follow potential changes in the Fe concentration



and Fe isotope composition of the soil surrounding the mesh bags.

Solid phase analysis

Solid samples from ^{57}Fe -mineral mesh bags without soil were analyzed by bulk Fe K-edge (7112 eV) X-ray absorption spectroscopy (XAS) at the SAMBA beamline at the SOLEIL synchrotron (St. Aubin, France), and by powder XRD. X-ray absorption near-edge structure and extended X-ray absorption fine structure (EXAFS) spectra were recorded as transmission spectra. Linear combination fit (LCF) analyses of k^3 -weighted EXAFS spectra were performed in Athena.⁴⁹ Reference compounds for Fe-minerals comprised ferrihydrite, lepidocrocite, magnetite, goethite, siderite, and green rust carbonate. Rietveld quantitative phase analysis (QPA)⁵⁰ of XRD patterns was used to obtain the mineral fractions of transformation products in the samples. Ferrihydrite was included in the QPA as a mass-calibrated PONKCS⁵¹ phase, as described in earlier studies.^{14,18,23} Detailed information on sample preparation, measurements, fitting of XAS spectra and Rietveld analyses are found in ESI Sections S7 and S8.†

The mineral transformation in ^{57}Fe -Fh- and ^{57}Fe -Lp-soil mixes during the incubation was investigated by ^{57}Fe Mössbauer spectroscopy, which is only sensitive to ^{57}Fe in the sample. With 8 mg of ^{57}Fe -labeled mineral (^{57}Fe fraction = 95.08%) mixed with 800 mg of soil (Fe = 3.3 g kg^{-1}) with ^{57}Fe isotope composition (2.12% ^{57}Fe ; ref. 52), >98% of the Mössbauer signal came from the introduced mineral Fe, while $\leq 2\%$ of the signal originated from the soil Fe. Additionally, ^{57}Fe -mineral samples collected at the end of anoxic (11 weeks) and oxic (12 weeks) periods of redox cycle III in the redox fluctuating mesocosms and at 12 weeks from the permanently flooded mesocosm were also analyzed with Mössbauer spectroscopy. The Mössbauer spectra were collected at 77 K and 5 K in transmission mode and fitted using the Recoil software.⁵³ The ^{57}Fe -mineral-soil mixes sampled after redox cycle III (12 weeks) were additionally measured at room temperature (295 K) to consider potential magnetite formation. Details to sample preparation and fitting of Mössbauer spectra are presented in ESI Section S11.†

Mineral dissolution and aqua regia digestions

The elemental composition of initial and incubated ^{57}Fe -mineral samples was determined after dissolution in concentrated HCl (NORMATOM®, VWR) at room temperature by ICP-OES. The C content of ^{57}Fe -Fh and ^{57}Fe -Lp at the end of the last redox cycle (12 weeks) was determined using a CHN analyzer (LECO TruSpec Micro). To measure changes in the Fe content and Fe isotope composition, initial and reacted ^{57}Fe -mineral-soil mixes and soil samples from Humax cores were digested with aqua regia. Aliquots (~150 mg) of the soil and/or mineral-soil mixes were weighed into 15 mL centrifuge tubes and 10 mL of freshly prepared aqua regia ($\text{HNO}_3:\text{HCl}$ ratio 1:3, HNO_3 (distilled; Merck), HCl (NORMATOM®, VWR)) was added to each vial. The samples were digested at 120 °C for 90 min and passed through a 0.45 μm PTFE filter. The total Fe concentration in the filtrates was

measured by ICP-OES. For Fe isotope analyses, all samples were diluted to ~50 ppb Fe and analyzed with triple-quadrupole inductively coupled plasma mass spectrometry (ICP-MS, Agilent 8800 Triple Quad), and ^{57}Fe isotope fractions ($f^{57}\text{Fe}$) were calculated relative to the sum (counts per second) of ^{54}Fe , ^{56}Fe , ^{57}Fe , and ^{58}Fe .^{14,23}

Results and discussion

Soil conditions during the experiment

Matric potentials (ψ) showed that the soil was water saturated during flooded periods ($\psi = 0$ kPa), while the water content dropped to slightly below field capacity during drained periods ($\psi = -9$ kPa, field capacity = -6 kPa; Fig. S8†). The pH dropped from circumneutral conditions (pH 6.5–6.9) during flooded periods to slightly acidic conditions (pH 5.7–6.0) during drained periods (Fig. S9†). The pH was slightly higher in the last, compared to the first flooded period. The soil Eh in the mesocosms with fluctuating water table indicated Fe reducing soil conditions (<100 mV at pH 7; ref. 9 and 54) during flooded periods (-163 ± 22 mV, mean Eh at 3, 7 and 11 weeks \pm standard error of the mean, Fig. S8†), while the Eh was well above conditions expected for Fe reduction during the drained periods ($+386 \pm 25$ mV, mean Eh at 4, 8 and 12 weeks). After anoxic conditions had established in the permanently flooded mesocosm, the Eh remained stable at -141 ± 7 mV (mean Eh at 2–12 weeks) throughout the experiment (Fig. S8†).

Porewater element concentrations

In the redox fluctuating mesocosms, Fe was released from the solid phase to the porewater within one day after soil flooding (Fig. 1A). The dissolved Fe concentration was highest one week after flooding (1.1 mM), followed by a slight decrease to 0.9 mM Fe at the end of the anoxic period of redox cycle I (3 weeks). In the following two redox cycles (II and III), dissolved Fe dynamics were similar to redox cycle I, but Fe release was slower and peak Fe concentrations were lower. Combined with measured changes in Eh and pH (Fig. S8 and S9†), the Fe release to the porewater indicates that solid phase Fe(III) was reductively dissolved during anoxic periods.

Dissolved organic C and dissolved Si and P were present in the porewater at one week (249 mg L^{-1} DOC, 0.3 mM Si, 0.1 mM P, Fig. 1B–D), along with major base cations (3.8 mM Ca, 8.0 mM Na; initial flooding water contained 1 mM Ca, 5 mM Na) and small amounts of K and Mg (Fig. S10†). Concentrations of other elements were below detection limit. Since major base cations, such as Ca and Mg, have a relatively low affinity for adsorption to ferrihydrite,⁵⁵ the direct interactions between Ca, Mg, K and Na with Fe oxyhydroxides were likely much weaker than for DOC⁵⁶ and negatively charged oxyanions, such as phosphate⁵⁷ and silicate.^{57,58} Therefore, here we will focus on the element dynamics of Fe, DOC, Si, and P. The concentrations of DOC, Si, and P showed trends similar to dissolved Fe during redox cycles, with low concentrations at the start of anoxic periods followed by an increase in concentrations in the following days. The similar trends in Fe compared to DOC, Si,





Fig. 1 Total dissolved concentrations of (A) iron (Fe), (B) organic carbon (DOC), (C) silicon (Si), and (D) phosphorus (P) in porewaters of soils in the redox fluctuating (RF 1–3) and the permanently flooded mesocosms. Labels above grey boxes indicate anoxic (anox, grey boxes) and oxic (ox) periods in redox cycles I–III. Error bars show the standard error of triplicate porewater samples in each mesocosm. Dashed lines are included for visual aid.

and P release during anoxic periods suggest that these components were associated with Fe minerals and released to solution upon Fe mineral reductive dissolution. Additional DOC, Si, and P may have been released from metabolized organic matter.

In the permanently flooded mesocosm, the element concentrations in the porewater in the first three weeks were similar to the concentrations observed during the anoxic period of redox cycle I in the redox fluctuating mesocosms (Fig. 1). Dissolved Fe and DOC concentrations reached their maximums at one week (0.9 mM Fe, 183 mg L⁻¹ DOC), followed by a steady decrease to 0.4 mM Fe and 78 mg L⁻¹ DOC at the end of the experiment (12 weeks, Fig. 1A and B). Si and P were released within one week after the flooding of the soil in the permanently flooded mesocosm (0.3 mM Si and 0.1 mM P at 1 week) and subsequently increased up to 0.5 mM Si and 0.15 mM P at the end of the experiment (12 weeks, Fig. 1C and D).

The dissolved Fe, DOC and P concentrations reached a similar level in the permanently flooded mesocosm compared

to the end of each anoxic period in the redox fluctuating mesocosms (Fig. 1A, B and D). This suggests that the soil replenished any C and P that was lost with the drainage water in the redox fluctuating mesocosms upon the next flooding. The dissolved Si concentrations at the end of the anoxic periods of redox cycles II and III were slightly lower compared to the permanently flooded mesocosm. This may be explained by interrupted silicate dissolution during oxic periods in the redox fluctuating mesocosms. Additionally, Si polymerization may have been enhanced due to lower pH and the relative increase in Si concentrations^{59,60} during oxic periods in the redox fluctuating mesocosms.

^{NA}Fe-mineral transformations in mesh bags without soil

The ^{NA}Fe-Fh and ^{NA}Fe-Lp in mesh bags without soil transformed to more crystalline Fe minerals over the three redox cycles, as shown by linear combination fitting of Fe K-edge EXAFS spectra (Fig. 2 and S15[†]). During the anoxic period of



redox cycle I, ^{54}Fe -Fh transformed to goethite (52% of total Fe) and magnetite (30%) (Fig. 2A). During the following oxic period, fractions of ferrihydrite increased in ^{54}Fe -Fh, while goethite fractions remained relatively stable and magnetite fractions decreased. At the end of redox cycle I (4 weeks), ^{54}Fe -Fh samples contained 33% ferrihydrite, 51% goethite and 16% magnetite. In comparison, ^{54}Fe -Lp transformed to magnetite (26%) during the anoxic period of redox cycle I (Fig. 2B) and the (nano-) goethite that was present in the initial ^{54}Fe -Lp (ESI Section S2†) disappeared. With the subsequent oxic period, fractions of lepidocrocite increased and magnetite fractions decreased, leading to 94% lepidocrocite and 6% magnetite in ^{54}Fe -Lp samples after redox cycle I (4 weeks). Similar trends in Fe mineral fractions were observed during redox cycles II and III, but changes in mineral fractions between anoxic and oxic periods were smaller.

The presence of goethite and/or magnetite in the incubated ^{54}Fe -Fh and ^{54}Fe -Lp samples was also confirmed by Mössbauer spectroscopy (Fig. 3 and Table 1) and XRD (Fig. S12 and S13†). Mössbauer spectra (5 K) of samples from redox cycle III showed sextets with parameters matching reported values of Fe in magnetite.⁶¹ Fitting of magnetite in Mössbauer spectra at low temperature is challenging since magnetite contributes up to five sextets at 5 K.⁶¹ We fit the spectra following the approach of Doriguetto *et al.*⁶¹ (Fig. S19A and B†). Since two of the five sextets covered $\geq 80\%$ of the sextet area attributed to magnetite, we continued with fitting only the two dominant sextets (Fig. S19C and D†). These sextets correspond to sextets S4 and S5 in Fig. 3 and in the following text. According to Doriguetto *et al.*,⁶¹ the parameters of sextets S4 and S5 represent octahedral $\text{Fe}^{2,67+}$ and tetrahedral Fe^{3+} in magnetite, respectively. In addition to Fe mineral phases detected by EXAFS and Mössbauer, XRD

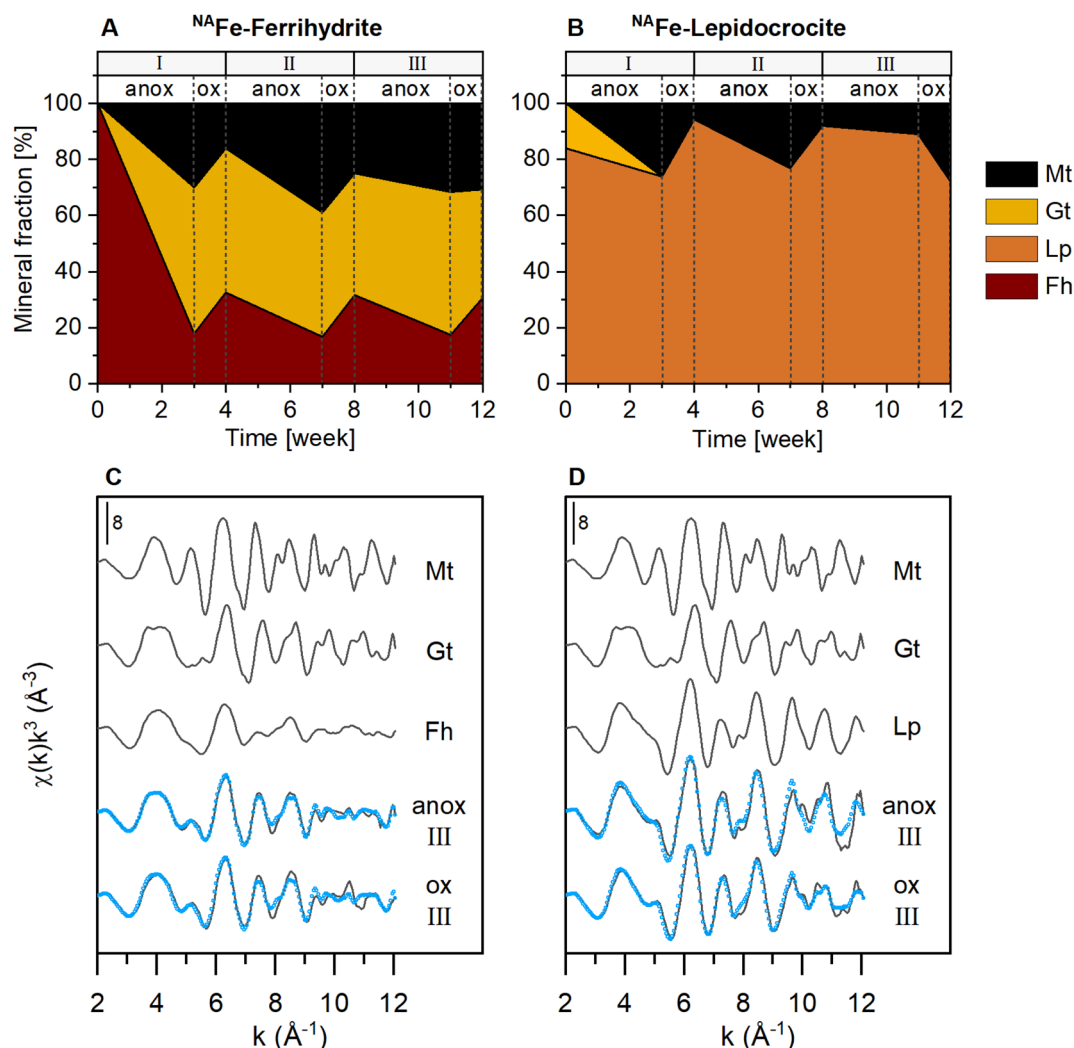


Fig. 2 Iron mineral fractions as determined by linear combination fitting of Fe K-edge EXAFS spectra in ^{54}Fe -mineral samples from mesh bags without soil for (A) ^{54}Fe -ferrihydrate and (B) ^{54}Fe -lepidocrocite incubated for 12 weeks in the redox fluctuating mesocosms, with (C and D) reference spectra and spectra from redox cycle III. According to EXAFS data, the initial ^{54}Fe -lepidocrocite contained minor fractions (16%) of goethite which was likely nano-crystalline (ESI Section S2†). All spectra and fits are presented in Fig. S15.† Fractions of fit components and reduced χ^2 are presented in Table S4.† Abbreviations: Mt = magnetite, Gt = goethite, Fh = ferrihydrite, Lp = lepidocrocite, anox = anoxic, ox = oxic.



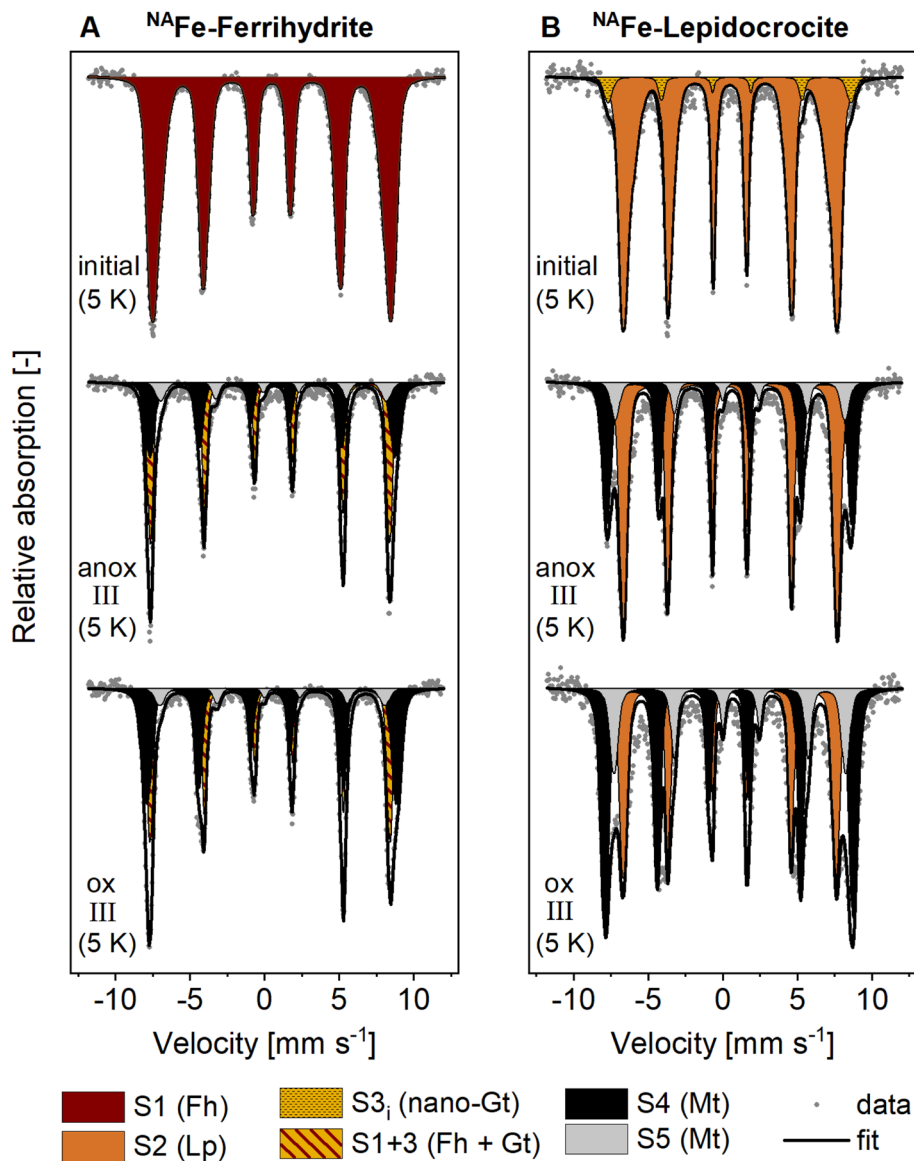


Fig. 3 Mössbauer spectra collected at 5 K from (A) ^{54}Fe -ferrihydrate and (B) ^{54}Fe -lepidocrocite mesh bags without soil before the incubation (initial) and after the incubation in the redox fluctuating mesocosms (redox cycle III anoxic and oxic period). Averaged fitting parameters and interpretations of Mössbauer fit components are given in Table 1, all fitting parameters are presented in Tables S11 and S12.† Abbreviations: anox = anoxic, ox = oxic, S = sextet.

identified small fractions ($\leq 5\%$) of siderite in ^{54}Fe -Fh and ^{54}Fe -Lp samples from anoxic and oxic periods (Fig. S12†). The applied methods for the identification and quantification of mineral phases (Fe K-edge EXAFS, Mössbauer spectroscopy, XRD) generally agreed. A comparison of quantified mineral fractions from all methods is presented for samples from redox cycle III and the permanently flooded mesocosm (12 weeks) in Table S13.†

The decrease in magnetite fractions observed in ^{54}Fe -Fh and ^{54}Fe -Lp during the oxic periods may be related to the oxidation of solid-associated Fe(II) and its precipitation as ferrihydrate, lepidocrocite or goethite. This may have led to a relative decrease in magnetite fractions. Magnetite also underwent changes in the Fe(II) : Fe(III) stoichiometry during redox cycles,

especially in ^{54}Fe -Fh samples. The Mössbauer fitting results indicate a decrease in the ratio between areas of sextets S4 (Fe(II) and Fe(III)) and S5 (only Fe(III)) during the oxic period (Fig. 3) which suggests that the Fe(II) : Fe(III) ratio in magnetite was higher during anoxic compared to oxic periods. This is supported by Rietveld fitting results from XRD patterns, which show a larger unit cell length (a) in magnetite in anoxic (8.381 Å, redox cycle I) compared to oxic (8.365 Å, redox cycle I) periods (Fig. S14†). Similar decreases in the magnetite unit cell length have previously been related to a decreased Fe(II) : Fe(III) ratio in magnetite⁶² (ESI Section S7†).

Since the ^{54}Fe -mineral samples were spatially separated from the soil, a large part of the mineral transformations were likely caused by the interactions with Fe(II) from the porewater



Table 1 Fit components, averaged fitting parameters and corresponding interpretations for Mössbauer spectra from ^{57}Fe -minerals and ^{57}Fe -mineral-soil mixes collected at 5 K. All fitting parameters are presented in ESI Section S11

Fit component	CS ^a [mm s ⁻¹]	QS ^b or ϵ ^c [mm s ⁻¹]	H^d [T]	Interpretation	Ref.
D1	0.47	0.84	—	Complexed and silicate-associated Fe(III)	35
D2	1.22	2.62	—	Adsorbed Fe(II) or Fe(II) in primary minerals or silicates	35 and 45
S1	0.47	< -0.01	48.84	Fe(III) in ferrihydrite	67
S2	0.48	0.01	42.68	Fe(III) in lepidocrocite	68
S3	0.48	-0.13	49.89	Fe(III) in goethite	45
S3 _i	0.51	-0.06	50.45	Nano-crystalline goethite	45
S1+3	0.48	-0.06	47.15	Mixture of ferrihydrite and goethite	—
S4	0.44	0.02	51.18	Octahedral Fe(II) and Fe(III) in magnetite	61
S5	0.85	-0.34	47.47	Tetrahedral Fe(III) in magnetite	61
CF	0.81	0.00	26.31	(Mixed-valence) disordered Fe phase	35, 37 and 45

^a CS: center shift. ^b QS: quadrupole splitting (for doublets). ^c ϵ : quadrupole shift (for sextets). ^d H : hyperfine field. Abbreviations: D = doublet, S = sextet, CF = collapsed feature.

diffusing into the mesh bags, as suggested by a recent study using confocal Raman microspectroscopy.³⁰ In turn, microbial reduction of ferrihydrite and lepidocrocite inside the mesh bags likely played only a minor role in mineral transformation. That Fe(II) catalyzed the transformation of ferrihydrite and lepidocrocite in ^{57}Fe -mineral mesh bags without soil is consistent with the fact that goethite is a frequent product of Fe(II)-catalyzed ferrihydrite transformations in model experiments with mineral suspensions spiked with aqueous Fe(II).^{12–17} The formation of goethite from ferrihydrite has also been observed in similar incubation studies using mineral mesh bags in flooded soils.^{30,31} However, goethite only formed from ^{57}Fe -Fh, but not from ^{57}Fe -Lp samples, whereas magnetite formed from both ^{57}Fe -Fh and ^{57}Fe -Lp. This observation agrees with recent reports from Fe(II)-catalyzed mineral transformation experiments, where goethite nucleated from ferrihydrite but not from lepidocrocite, while magnetite formed from both minerals.^{14,63} Considering the slightly acidic conditions of the soil in the current study (Fig. S9†) as well as the presence of dissolved (Fig. 1) and sorbed (Fig. S11†) C, Si and P, the transformation to magnetite was somewhat unexpected. Magnetite has been reported as a product of Fe(II)-catalyzed ferrihydrite and lepidocrocite transformation at higher pH (≥ 7)^{12,18} and the presence of organic matter or Si can hinder the formation of magnetite.^{14,23} However, solid concentrations of C, Si and P were low (molar ratios of C/Fe; up to 0.1%, Si/Fe; up to 1.3%, and P/Fe; up to 0.9%) compared to element ratios reported in other studies in which magnetite formed (e.g., C/Fe ratio of 0.7 (ref. 23) and P/Fe ratios of ~ 0.05 (estimated from the reported value of 646 mmol P kg⁻¹ ferrihydrite)⁶⁴). Additionally, C, Si, and P may have accumulated in the outermost mineral layer of the ^{57}Fe -Fh and ^{57}Fe -Lp mesh bags,³⁰ as the interaction of ^{57}Fe -Fh and ^{57}Fe -Lp with dissolved elements in the porewater was controlled by the diffusion of these elements into the mineral mesh bags. However, despite the accumulation of sorbed elements, electron transfer between dissolved Fe(II) and structural Fe(III) can still occur.^{14,23,65} This likely enabled mineral transformation to crystalline Fe minerals inside the mesh

bags.³⁰ Magnetite formation during anoxic periods would have been supported by locally high Fe(II) concentrations. Locally high Fe(II) concentrations may have been established due to a Fe(II) diffusion gradient towards the mineral mesh bags, caused by the large sorption capacity of ferrihydrite and lepidocrocite for Fe(II) in mineral mesh bags. Considering the similar use of mineral mesh bags, it is intriguing that Grigg *et al.*³⁰ observed goethite but no magnetite formation. A potential reason for the formation of magnetite in the present study is the 5-times lower amount of ferrihydrite in the mesh bags. Mesh bags with a lower amount of ferrihydrite contain less Fe(III) and thus require smaller amounts of Fe(II) to achieve sufficiently high Fe(II):Fe(III) ratios for magnetite formation. Once magnetite nucleates, its accumulation quickly captures additional Fe(II) from solution.^{14,21,66}

Interpretation of Mössbauer fit components for ^{57}Fe -mineral-soil mixes

In contrast to the ^{57}Fe -mineral mesh bags without soil, the synthetic ferrihydrite and lepidocrocite in the ^{57}Fe -mineral-soil mixes constituted only a small fraction (1% w/w) of the samples (8 mg of ^{57}Fe -Fh or ^{57}Fe -Lp added to 800 mg of soil). Therefore, XRD could not be used to follow the transformations of the added minerals. Instead, we used Mössbauer spectroscopy, which selectively detects ^{57}Fe in the samples, of which >98% originated from the added ^{57}Fe -labeled minerals. In the initial ^{57}Fe -Fh-soil mix (Fig. 4C), the fitting parameters of the sextet (center shift, CS = 0.47 mm s⁻¹; quadrupole shift, ϵ = -0.007 mm s⁻¹; hyperfine field, H = 48.84 T), matched with parameters reported for ferrihydrite.⁶⁷ Since fitting ferrihydrite in Mössbauer spectra is challenging⁶⁷ and our samples also contain multiple other sextets at 5 K, we decided to fix the ferrihydrite parameters, as derived from the initial ^{57}Fe -Fh-soil mix, for subsequent fits. The initial ^{57}Fe -Lp-soil mix (Fig. 4D) showed a sextet with a CS of 0.49 mm s⁻¹, an ϵ of 0.03 mm s⁻¹ and a H of 44.35 T, which agrees with parameters reported for lepidocrocite.⁶⁸ In the initial ^{57}Fe -Lp, a second sextet (S3_i) with parameters (CS = 0.49 mm s⁻¹, ϵ = -0.05 mm s⁻¹, H = 50.37 T),





Fig. 4 Spectral area of fit components in 5 K Mössbauer spectra for (A) ^{57}Fe -ferrihydrate-soil mixes, and (B) ^{57}Fe -lepidocrocite-soil mixes incubated in the redox fluctuating mesocosms. Lower panels show Mössbauer spectra collected at 5 K from initial mineral-soil mixes containing (C) ^{57}Fe -ferrihydrate or (D) ^{57}Fe -lepidocrocite and corresponding samples after incubation in the redox fluctuating mesocosms, sampled in the anoxic (11 weeks) and oxic (12 weeks) period of redox cycle III. Abbreviations: anox = anoxic, ox = oxic, D = doublet, S = sextet, CF = collapsed feature, Fh = ferrihydrate, Lp = lepidocrocite, Gt = goethite. Goethite in initial ^{57}Fe -lepidocrocite likely was nano-crystalline. Interpretations of Mössbauer fit components are given in Table 1, fitting parameters are presented in Tables S9 and S10.†

indicated that nano-goethite was present in this sample (more details are presented in ESI Section S2†).

The fit components for Mössbauer spectra of the incubated ^{57}Fe -Fh and ^{57}Fe -Lp-soil mixes collected at 5 K, averaged fitting parameters and corresponding interpretations are summarized in Table 1. Results from 77 K spectra are presented in Fig. S20.† All fitted parameters are reported in Tables S7–S10.† The Fe(III) doublet (D1) parameters at 5 K are similar to parameters

indicating pyrite.⁶⁹ However, pyrite formation is unlikely since total dissolved S concentrations in this experiment were below the detection limit. Therefore, in this study the Fe(III) doublet (D1) is interpreted as complexed and silicate-associated Fe(III) , in agreement with previous studies.^{35,45} The fitting of 5 K spectra collected from ^{57}Fe -Fh and ^{57}Fe -Lp-soil mixes required a collapsed feature. In this work, we interpreted the collapsed feature as a highly disordered Fe phase,^{35,37,45} which in our case



was a mixed-valence Fe(II)–Fe(III) phase with large fractions of Fe(II), as seen from differences in Fe(II) fractions in 77 K and 5 K spectra (Fig. S23†). A detailed discussion on the nature of the collapsed feature, including results from sequential Fe extractions (ESI Section S10†), and fitting results from 77 K spectra (ESI Section S11†) are found in the ESI.†

Over the experiment duration, we observed increased noise in Mössbauer spectra from incubated ^{57}Fe -mineral-soil mixes (compare spectra of initial and incubated mineral-soil mixes in Fig. 4C and D). The results from aqua regia digestions of the ^{57}Fe -mineral-soil mixes showed decreasing Fe contents and decreasing ^{57}Fe fractions over time (Fig. S16A and B†). Based on this, we infer that the increased spectral noise was due to the reductive dissolution of ^{57}Fe -minerals and the loss of ^{57}Fe (II) from the mesh bags through diffusion and water flow during drainage. This is supported by the higher loss of ^{57}Fe from ^{57}Fe -Fh compared to ^{57}Fe -Lp-soil mixes, which agrees with the lower stability of ferrihydrite during reductive dissolution.²⁰ Despite the increased spectral noise, the fits were robust, and we do not expect a major change in the interpretation of the Mössbauer spectra. Regardless of the loss of ^{57}Fe from the mesh bags, the majority of the Mössbauer signal still originated from the added ^{57}Fe -labeled ferrihydrite (>71%) or lepidocrocite (>84%, Fig. S16C†). Nevertheless, the identification and quantification of minor fit components (<10% of fitted spectral area) should be interpreted with caution due to the spectral noise. The stated ^{57}Fe phase fractions in the following text are derived from the fitting of Mössbauer spectra and thus represent relative fractions of the ^{57}Fe that remained in the mesh bags. The results from aqua regia digestions of the initial and incubated ^{57}Fe -mineral-soil mixes and the surrounding soil are presented in ESI Section S9.†

^{57}Fe -mineral transformations in mesh bags with soil

Mineral transformations in ^{57}Fe -Fh-soil mixes resulted in less crystalline mineral transformation products and a larger variety of Fe phases compared to ^{57}Fe -Fh samples without soil (compare Fig. 3 with Fig. 4). During the anoxic period of redox cycle I (3 weeks), the fraction of ferrihydrite in the ^{57}Fe -Fh-soil mix decreased from 100% to 37% of total ^{57}Fe , while the fraction of lepidocrocite (24%) and the mixed-valence disordered Fe phase (26%) increased (Fig. 4A). Additionally, small fractions (each $\leq 8\%$) of adsorbed/clay-associated Fe(II) and organically-complexed/silicate Fe(III) formed. Upon the drainage of the soil, the fraction of lepidocrocite (32%) increased and the fraction of the mixed-valence disordered Fe phase (19%) decreased. The fraction of ferrihydrite remained relatively stable (39%). Overall, these results indicate that the ^{57}Fe -Fh was reductively dissolved and partially transformed to lepidocrocite and the mixed-valence disordered Fe phase during the first anoxic period. During the subsequent oxic period, part of the adsorbed Fe(II) and Fe(II) in the mixed-valence disordered Fe phase was oxidized and precipitated as lepidocrocite or ferrihydrite. The formation of lepidocrocite and the mixed-valence disordered Fe phase in the ^{57}Fe -Fh-soil mixes contrasts the results from ^{57}Fe -Fh samples without soil, where goethite and magnetite formed.

In the following redox cycles (II and III), the fractions of Fe(II)-bearing phases (adsorbed/clay-associated Fe(II), mixed-valence disordered Fe phase) in the ^{57}Fe -Fh-soil mixes increased during anoxic and decreased during oxic periods. Over the three redox cycles, the contribution of Fe(II)-bearing phases in ^{57}Fe -Fh-soil mixes generally increased (*cf.* fractions of Fe(II) and the mixed-valence disordered Fe phase in anoxic periods of redox cycles I–III, Fig. 4A). As opposed to trends in Fe(II)-bearing phases, the fractions of Fe(III)-phases (ferrihydrite, lepidocrocite, organically-complexed/silicate-associated Fe(III)) decreased during anoxic and increased during oxic periods, with most pronounced changes for lepidocrocite fractions. Lepidocrocite in the ^{57}Fe -Fh-soil mixes disappeared quickly during the anoxic periods in redox cycles II and III, indicating that the newly formed lepidocrocite was highly susceptible to reductive dissolution. The fast reductive dissolution of lepidocrocite was likely promoted by coprecipitation with other dissolved components in the porewater, which can lead to a decreased crystallinity of lepidocrocite.^{70,71}

In ^{57}Fe -Lp-soil mixes, the lepidocrocite fraction decreased from 84% to 25% during the anoxic period of redox cycle I (3 weeks, Fig. 4B). In contrast to magnetite formation in ^{57}Fe -Lp mesh bags without soil, the main mineral transformation products in the ^{57}Fe -Lp-soil mixes were ferrihydrite (35%) and goethite (25%). The nano-goethite that was detected in initial ^{57}Fe -Lp (ESI Section S2†) was not found in the incubated ^{57}Fe -Lp-soil mixes. Also, small contributions (each $\leq 7\%$) from adsorbed/clay-associated Fe(II), organically-complexed/silicate-associated Fe(III) and the mixed-valence disordered Fe phase were present. Upon the soil drainage in redox cycle I, the oxidation state and speciation of ^{57}Fe atoms in ^{57}Fe -Lp-soil mixes were relatively stable. However, in the following redox cycles (II and III), fractions of the Fe(II)-bearing phases (adsorbed/clay-associated Fe(II), mixed-valence disordered Fe phase) increased slightly during anoxic periods and decreased during oxic periods, but changes were smaller compared to the ^{57}Fe -Fh-soil mixes. The contribution of lepidocrocite in ^{57}Fe -Lp-soil mixes generally decreased over the redox cycles and had almost completely disappeared ($\leq 6\%$) at the end of redox cycle III (12 weeks, Fig. 4D). In contrast to lepidocrocite fractions, the goethite fraction increased during anoxic periods and remained stable during oxic periods. This led to a general increase of the goethite fraction over the redox cycles, with goethite accounting for up to 45% of ^{57}Fe at the end of redox cycle III (12 weeks). This suggests that lepidocrocite and ferrihydrite in the ^{57}Fe -Lp-soil mixes were preferentially reductively dissolved during anoxic periods, promoting the formation of more goethite. Goethite present in the soil or nano-goethite in initial ^{57}Fe -Lp may have additionally supported the formation of goethite in these samples.^{63,72} In contrast to magnetite accumulation in the ^{57}Fe -mineral mesh bags without soil, no indication of magnetite was found in the ^{57}Fe -mineral-soil mixes. The absence of magnetite in incubated ^{57}Fe -mineral-soil mixes was further tested by collecting room temperature Mössbauer spectra, where magnetite would form two sextets.^{64,73} No sextet was present in the room temperature Mössbauer spectra of ^{57}Fe -Fh and ^{57}Fe -Lp-soil mixes (redox cycle III, oxic, Fig. S22†).



The fraction of the mixed-valence disordered Fe phase increased during anoxic periods in both the ^{57}Fe -Fh and ^{57}Fe -Lp-soil mixes. During oxic periods, most of the Fe(II) in the mixed-valence disordered Fe phase was oxidized and its fraction decreased. Especially in ^{57}Fe -Fh-soil mixes, fractions of the mixed-valence disordered Fe phase showed drastic changes between anoxic and oxic periods (Fig. 4A). The formation of the mixed-valence disordered Fe phase in the ^{57}Fe -mineral-soil mixes during anoxic periods was likely related to the close association of Fe minerals with the soil matrix. This likely hindered the recrystallization of ferrihydrite and lepidocrocite through the direct exposure to, for example, DOC and dissolved Si and P. Assuming a dissolution-precipitation mechanism for mineral recrystallization,^{63,74,75} these findings agree with the hindered crystallization of Fe oxyhydroxides in the presence of organic matter,⁷⁶ Si,^{77–79} and P.^{57,71,78} Hindered mineral recrystallization likely also supported ferrihydrite formation in ^{57}Fe -Lp samples. Similarly, a recent study that incubated ^{57}Fe -labeled ferrihydrite in soil microcosms reported ferrihydrite transformation to a green rust-like phase with no sign of lepidocrocite or goethite formation.³³ Despite the use of a similar soil as in the study of Notini *et al.*,³³ in the present study, no green rust was found, which may be related to the lower pH⁸⁰ (pH 6.9 in this study, compared to pH 7.6 in Notini *et al.*³³). However, we cannot exclude small contributions of green rust in the collapsed feature observed in Mössbauer spectra.

The changes in solid-phase ^{57}Fe speciation in the ^{57}Fe -Fh and ^{57}Fe -Lp-soil mixes induced by the onset of anoxic and oxic soil conditions became larger with every redox cycle. For example, the fractions of Fe(II)-bearing phases (adsorbed/clay-associated Fe(II), mixed-valence disordered Fe phase) during anoxic periods increased with every redox cycle (Fig. 4A and B). The increasingly reduced state of solid-associated ^{57}Fe in ^{57}Fe -Fh and ^{57}Fe -Lp samples after incubation observed in 5 K Mössbauer spectra was also supported by 77 K spectra collected from anoxic samples. In 77 K spectra, the fraction of a doublet ($CS = 1.29 \text{ mm s}^{-1}$; quadrupole splitting, $QS = 2.62 \text{ mm s}^{-1}$), which corresponds to Fe(II),⁴⁵ increased with every redox cycle up to 76% and up to 32% of total ^{57}Fe in ^{57}Fe -Fh- and ^{57}Fe -Lp-soil mixes, respectively (redox cycle III, anoxic period, Fig. S20†). This can likely be explained by the precipitation of more easily reducible Fe phases during the oxic phases, facilitating faster microbial Fe reduction during the subsequent anoxic periods.^{39,81} The increasing fractions of Fe(II)-bearing phases and the increasingly pronounced changes in the ^{57}Fe oxidation state demonstrate that the susceptibility of Fe phases to reductive dissolution increased with repeated soil redox cycles.

Studies using soils from contrasting locations found that fractions of short-range-ordered Fe phases can increase^{34,35,37} or decrease^{35,37,38} during soil redox fluctuations. For example, Thompson *et al.*³⁸ studied highly weathered tropical forest soils, where Fe mineralogy was dominated by nano- and micro-crystalline goethite (determined by Mössbauer spectroscopy) and found increasing Fe crystallinity after multiple complete redox cycles (4 cycles within 8 weeks). In comparison, Thomas-Arrigo and Kretzschmar³⁶ investigated organic matter-rich soils from Iceland, where ferrihydrite, goethite, and some

lepidocrocite were the main Fe mineral phases (determined by Fe K-edge XAS) and reported similar Fe mineral composition before and after the exposure to one complete redox cycle with varying duration (1–5 weeks anoxic, 2–8 days oxic). These studies shed light on the redox transformation of native soil minerals, which included multiple Fe phases. With the use of ^{57}Fe -labeled minerals, the present study particularly demonstrated the transformation of ferrihydrite and lepidocrocite during three redox cycles in a sandy and weakly acidic paddy soil which was poor in organic matter. The outcomes of this study suggest that the stability of Fe oxyhydroxides during mineral transformation and reductive dissolution strongly impact mineral transformations towards lower or higher crystallinity during soil redox fluctuations.

Mineral transformations in the permanently flooded mesocosm

For both the ^{57}Fe -mineral samples (Fig. 5A) and the ^{57}Fe -mineral-soil mixes (Fig. 5B), the same mineral phases formed from minerals incubated in the permanently flooded (12 weeks) and the redox fluctuating mesocosms (redox cycle III, anoxic period). However, the relative fractions of Fe phases varied. The ^{57}Fe -Fh in the permanently flooded mesocosm transformed to similar amounts of goethite (48%) but more magnetite (46%) compared to ^{57}Fe -Fh in the redox fluctuating mesocosms (51% goethite, 32% magnetite, Fig. 5A). Thus, the transformation extent of ^{57}Fe -Fh was higher when incubated under permanently flooded conditions. Similarly, for ^{57}Fe -Lp, the incubation in the permanently flooded mesocosm resulted in a larger extent of lepidocrocite transformation and higher magnetite fractions (30%) compared to the redox fluctuating mesocosms (11% magnetite).

For ^{57}Fe -mineral-soil mixes, Mössbauer spectroscopy showed that in the ^{57}Fe -Fh-soil mixes, the ferrihydrite fraction was slightly lower after the incubation in the permanently flooded mesocosm (10%, 12 weeks) compared to the redox fluctuating mesocosms (23%, redox cycle III, anoxic period, Fig. 5B). In turn, the solid-associated Fe(II) fraction was slightly higher after the incubation under permanently flooded conditions (36% in permanently flooded, 28% in redox fluctuating mesocosms). This may reflect the regular precipitation of short-range-ordered Fe phases during oxic periods in the redox fluctuating mesocosms. In ^{57}Fe -Lp-soil mixes, slightly less goethite formed when incubated under permanently flooded conditions (34%) compared to redox fluctuating conditions (45%). Thus, the intermittent oxic periods may have supported goethite formation in ^{57}Fe -Lp-soil mixes.

The differences in overall mineral transformations in ^{57}Fe -mineral-soil mixes after the incubation in the permanently flooded or the redox fluctuating mesocosms were smaller than in ^{57}Fe -mineral mesh bags without soil. The higher transformation extent in ^{57}Fe -mineral mesh bags without soil incubated in the permanently flooded mesocosm is likely related to the continuous supply of Fe(II) from the Fe reducing soil matrix. In contrast, the Fe(II) supply to the ^{57}Fe -minerals in the mesh bags in the redox fluctuating mesocosms was likely interrupted during oxic





Fig. 5 Comparison between samples incubated in the redox fluctuating (RF, anoxic period of redox cycle III) and permanently flooded (PF, 12 weeks) mesocosms with fractions of fitted components in (A) Fe K-edge EXAFS spectra collected from ^{57}Fe -minerals and (B) Mössbauer spectra collected from ^{57}Fe -mineral-soil mixes at 5 K, with results for ferrihydrite (Fh) and lepidocrocite (Lp). Abbreviations: S = sextet, D = doublet, CF = collapsed feature, Gt = goethite, Mt = magnetite. Interpretation of Mössbauer fit components for panel B are presented in Table 1.

drained periods, stalling mineral transformations. Since the minerals in the ^{57}Fe -mineral-soil mixes were exposed to *in situ* reductive dissolution, the short-range-ordered Fe phases that formed during oxic periods likely were quickly reductively dissolved upon the onset of Fe reducing conditions. Accordingly, the drastic shifts in Fe crystallinity that occurred during oxic periods were largely reversed during the following anoxic periods. This allowed the progression of mineral transformations to continue as though the oxic interruption had not occurred.

The effect of close association with the soil on mineral transformations

Iron oxyhydroxide dynamics in soils are controlled by a balance between microbial reductive dissolution and mineral transformation or recrystallization, which determines the fate of the mineral. Our study demonstrates how this balance is impacted by the close association between minerals and the soil matrix and by initial mineral crystallinity. During the incubation of ^{57}Fe -mineral mesh bags without soil, magnetite and/or goethite were formed (Fig. 2 and 3). In contrast, when Fe minerals were closely associated with the soil in ^{57}Fe -mineral-soil mixes, minerals were reductively dissolved while mineral transformation was hindered. The hindered mineral transformation resulted in the formation of a mixed-valence disordered Fe phase (Fig. 4), the formation of which was promoted by fast reductive dissolution. This is supported by the fact that large fractions of the mixed-valence disordered Fe phase formed from poorly crystalline ferrihydrite in the ^{57}Fe -Fh-soil mixes (up to 46%, Fig. 4A). The higher stability of lepidocrocite during reductive dissolution²⁰ likely led to slower mineral dissolution in ^{57}Fe -Lp-soil mixes. Thus, the accumulation of the mixed-

valence disordered Fe phase was lower (up to 22%) and goethite formed (Fig. 4B). It is likely that the minerals in close contact with soil in the mineral-soil mixes were exposed to geochemical conditions more similar to those measured in the pore water outside the mesh bags compared to the geochemical conditions within the pure mineral mesh bags. As such, it is likely that the mineral transformations in the mineral soil mixes were strongly influenced by the presence of dissolved DOC, Fe and P, which ultimately hindered the transformation to crystalline mineral products. The formation of the mixed-valence disordered Fe phase in ^{57}Fe -mineral-soil mixes suggests that similar highly disordered and reactive Fe phases may also occur in natural soils.⁸²

Conclusions

This study investigated the transformation of Fe(III) oxyhydroxides during three soil redox cycles and the role of close contact between minerals and the soil. The transformation and recrystallization of ^{57}Fe -minerals in mesh bags without soil were likely mainly catalyzed by Fe(II) from the surrounding soil diffusing into the mesh bags and resulted in goethite and/or magnetite formation. Since Fe minerals in soils are part of the complex soil matrix, the results from ^{57}Fe -mineral-soil mixes may be more environmentally relevant. The presented findings highlight the complex interplay of biogeochemical factors directing Fe mineral transformations in redox-active soils. During the incubation as mineral-soil mixes, ferrihydrite was less stable during mineral transformation and reductive dissolution, compared to lepidocrocite. Thus, ferrihydrite was more strongly affected by redox cycles. Further,



mixing ferrihydrite with soil hindered mineral transformation compared to ferrihydrite samples that were not mixed with soil. For lepidocrocite, the close association with the soil enabled, and potentially even promoted, the transformation of lepidocrocite to goethite. Further, the close association of minerals with the soil promoted the formation of a highly disordered mixed-valence Fe(II)–Fe(III) phase. This was likely through a combination of *in situ* Fe reduction, Fe(II)-catalyzed mineral recrystallization and interactions with other dissolved soil components. It is possible that such highly disordered Fe phases serve as highly reactive sorbents for trace elements in paddy soils, impacting the availability of these elements for the uptake by the rice plants.

With repeated redox cycles, the production of solid-associated Fe(II) during anoxic periods increased likely due to the recurring formation of easily reducible short-range-ordered Fe phases during oxic periods. This highlights how the susceptibility of redox-affected soils to Fe reduction is influenced by the crystallinity of Fe phases in the soil and if or how frequently the soil has undergone redox fluctuations recently. Despite drastic changes in Fe crystallinity between anoxic and oxic periods, the redox cycling itself may only have a small impact on overall mineral transformations compared with permanent flooding of the same soil. In this respect, also the activity and composition of microbial communities and the availability of carbon sources need to be considered. Regarding coupled element cycles, the recurring formation of short-range-ordered Fe phases during redox cycles can provide a large sorption potential for trace elements. However, with repeated redox cycles the increased susceptibility of Fe to reductive dissolution during anoxic periods comes with an increasing risk of releasing mineral-associated elements to the porewater. Thus, the duration and frequency of flooded periods in paddy fields may impact the abundance of short-range-ordered Fe oxyhydroxides, their stability under Fe-reducing conditions and their potential to release associated trace-elements. These outcomes contribute to an improved understanding of Fe dynamics and potential implications for coupled element cycles during soil redox cycles.

Author contributions

Katrin Schulz: conceptualization, methodology, validation, investigation, formal analysis, writing – original draft, visualization; Luiza Notini: formal analysis, writing – review & editing; Andrew R. C. Grigg: methodology, writing – review & editing; L. Joëlle Kubeneck: investigation, writing – review & editing; Worachart Wisawapipat: methodology, writing – review & editing; Laurel K. ThomasArrigo: conceptualization, methodology, writing – review & editing, supervision; Ruben Kretzschmar: conceptualization, methodology, writing – review & editing, supervision, project administration, funding acquisition.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

We are grateful to Kurt Barmettler (ETH Zürich) for assisting with the soil sampling and laboratory analyses. We further thank the staff of the Ubon Ratchathani Rice Research Center for their support during the soil sampling, and the NRCT for granting the research permit (permit no. 0002/1164). Our thanks also goes to SOL-CONSEIL (Gland, France) for conducting the soil texture analysis, and to the MoBIAS laboratory (ETH Zürich, Switzerland), especially to Peter Kälin, for the analysis of carbon contents in ^{NA}Fe-mineral samples. We acknowledge SOLEIL (proposal no. 20221355) for the provision of synchrotron radiation facilities and thank Gautier Landrot (SAMBA) for his support during the synchrotron measurements. This work received funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme (grant agreement no. 788009-IRMIDYN-ERC-2017-ADG).

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