Isotopic variation of dissolved and colloidal iron and copper in a carbonatic floodplain soil after experimental flooding

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Abstract

Many floodplain soils worldwide are contaminated by present and past industrial and mining activities. During flooding redox potential decreases, triggering the release of dissolved and colloidal metals. We used an anaerobic microcosm incubation to simulate flooding of a carbonate-rich floodplain soil for 40 days. Soil solution samples were extracted to determine the release of dissolved (<0.02 µm) and colloidal fractions (0.02 - 10 µm). We analyzed stable isotope ratios of colloidal and dissolved Fe and Cu representing two groups of metals with different release behavior; release of Fe was steadily increasing, while Cu peaked sharply after flooding and decreased afterwards. The temporal trend of $\delta^{56}$Fe values of total Fe in
solution indicated dissimilatory iron reduction. The apparent isotopic fractionation between
dissolved and colloidal Fe \((\Delta^{56}\text{Fe}_{\text{dissolved-colloidal}} = \delta^{56}\text{Fe}_{\text{dissolved}} - \delta^{56}\text{Fe}_{\text{colloidal}})\) varied between
0.31±0.04‰ and -1.86±0.26‰. Low \(\delta^{56}\text{Fe}_{\text{colloidal}}\) (-1.16±0.04 ‰) values on day 4 of the
experiment suggested colloid formation by precipitation of dissolved Fe, while the strong
temporal variation in \(\Delta^{56}\text{Fe}_{\text{dissolved-colloidal}}\) indicated subsequent changes in colloid mineralogy,
sorption to soil components and/or electron transfer-atom exchange. The variations in \(\delta^{65}\text{Cu}\)
values \((\Delta^{65}\text{Cu}_{\text{dissolved-colloidal}}\) from 0.81±0.03‰ to 1.58±0.09‰) are probably linked to
changing oxidation state of colloidal Cu. While at the beginning of the experiment colloidal
Cu and solid soil Cu exchange, these systems decouple after the onset of sulfate reduction in
the second half of the experiment. The experimental results fit well to findings from
redoximorphic soils described in the literature and highlight the importance of colloids for
metal release and the isotopic pattern in carbonatic soils.

Keywords: Copper, Iron, Soil, Redox Processes, Colloids, Stable Metal Isotopes

Highlights

- Colloidal Fe precipitates from dissolved Fe in soil solution and does not originate from
  the mobilization of solid Fe minerals.
- The \(\delta^{56}\text{Fe}\) value of colloidal Fe changes with time indicating changes in mineralogy or
  electron transfer-atom exchange.
- Colloidal and solid soil Cu exchange occurs until sulfate reduction starts, afterwards
  colloidal Cu decouples from solid soil and colloids aggregate.
- The \(\Delta^{65}\text{Cu}_{\text{dissolved-colloidal}}\) values are probably driven by the redox state of colloidal Cu.
- Experimental findings agree well with field observations.
Graphical Abstract
1. Introduction

Present and past industrial and mining activities release contaminants into rivers, which continuously accumulate in floodplain soils (Burton et al., 2008; Frohne et al., 2011; Hindersmann and Mansfeldt, 2014). Depending on the frequency of flooding, metals might be remobilized and contaminate ground and surface waters (Du Laing et al., 2007; Hindersmann et al., 2014; Mansfeldt and Overesch, 2013). The mobility of metals mainly depends on organic matter concentrations and composition, soil mineral composition, redox conditions, and carbonate concentrations (Du Laing et al., 2009). When soils are flooded, the redox potential \(E_h\) decreases and Fe and Mn (oxyhydr)oxides are dissolved in many soils releasing associated trace elements (e.g., As, Ba, Co, Cr, V; Abgottspon et al., 2015; Hindersmann and Mansfeldt, 2014; Weber et al., 2009b). In some soils with permanent water saturation, the pool of reducible Fe and Mn is depleted and no further mobilization occurs under anoxic conditions. The release kinetics depends strongly on crystallinity of Fe and Mn minerals and microbial community composition in the soil (Lovley, 1991; Schuth et al., 2015). In carbonatic soils, the pH is buffered by the carbonic acid-hydrogen carbonate buffer system and proton release during formation of minerals like siderite and rhodochrosite (Abgottspon et al., 2015; Ponnamperuma, 1972). Furthermore, changes to anoxic conditions may cause microbial formation of reduced metal colloids (e.g., Cu\(^{0}\), Weber et al., 2009b). When \(E_h\) decreases, microbial sulfate reduction may be initialized and mobility of metals can be limited by the formation of or co-precipitation with sulfides (Borch et al., 2010; Weber et al., 2009a). However, sulfate reduction may also favor the release of metal-sulfide colloids of some elements (e.g., Ag, Cd, Cu, Hg, Pb) resulting in enhanced metal mobility during several days after flooding (Abgottspon et al., 2015; Hofacker et al., 2013; Weber et al., 2009b). The formation of sulfides is limited by the available sulfate content and controlled by the solubility of the corresponding metal sulfides (Weber et al., 2009a). Beside the formation of
organic, biomineralized and sulfide colloids, also colloids of Fe and Mn minerals may favor trace element transport (Hassellov and von der Kammer, 2008). The colloids may form because of precipitation of siderite and rhodochrosite in carbonatic soils or the precipitation of mixed valence (oxyhydr)oxides (e.g., magnetite or green rust; Ratering and Schnell, 2000; Tadanier et al., 2005; Taylor, 1980). Another colloid formation process is the partial reduction of ferrihydrite by microorganisms which may initialize deflocculation and release of the remaining ferrihydrite as colloids along with associated non-Fe elements (Tadanier et al., 2005). When soil flooding ends and redox conditions change back from anoxic to oxic, sulfides will be oxidized, releasing metals like Cd, Cu and Zn to pore waters, which may (co)precipitate with (oxyhydr)oxides (Contin et al., 2007; Du Laing et al., 2007; Frohne et al., 2011).

Abgottspon et al. (2015) differentiated trace metals into two groups with different release behavior in water-saturated soils. When $E_{h}$ decreased (from 300 mV to 0 mV), Group I metals (Ba, Co, Cr, Ni, and V) were constantly released during 40 days, predominantly in dissolved form (<50 % as colloids) closely correlated with the release of Fe and Mn by reductive dissolution of (oxyhydr)oxides. The increase in colloidal concentrations of Group I metals is probably attributable to coprecipitation of dissolved metals with and sorption to colloidal Fe and Mn minerals. Group II metals (Ag, Cd, Cu and Pb) were rapidly released after 2-6 days of flooding, strongly dominated by the colloidal fraction (>80%; 0.02-10 µm). In this study, we focus on the stable isotope ratios of Fe and Cu as representatives of Group I and Group II metals, respectively, to further explore the mechanisms underlying the observed differences in the release behavior of the two metal groups.

The stable isotope ratios of Fe and Cu bear a high potential to investigate long-term biogeochemical processes in soils, especially redox processes (Babcsanyi et al., 2014; Bigalke et al., 2013; Bigalke et al., 2010b; Mansfeldt et al., 2012; Wiederhold et al., 2007a). Iron is an important driver of biogeochemical processes and trace metal behavior in soils. Changing the
oxidation state of Fe results in marked Fe isotope fractionation causing variations in Fe isotopic composition among Fe pools in anoxic soils with δ⁵⁶Fe values ranging from -5‰ to +1‰ (Johnson et al., 2008). Under anoxic conditions, dissimilatory iron reduction (DIR) by bacteria initializes three main Fe reaction pools, i.e. Fe²⁺(aq), Fe²⁺(sorb) and Fe³⁺(reac). Some of the dissolved Fe²⁺(aq) may sorb to the oxide surface as Fe²⁺(sorb). Electron transfer between Fe²⁺(sorb) and Fe³⁺ creates a reactive layer of Fe³⁺ at the oxide surface (Fe³⁺(reac)) and is associated with ⁵⁶Fe enrichment in the solid phase. Subsequently, DIR releases isotopically light Fe into solution while heavy Fe isotopes are preferentially adsorbed on Fe oxide surfaces (Δ⁵⁶Fe_mineral-solution up to 4‰; Frierdich et al., 2014; Icopini et al., 2004; Mikutta et al., 2009; Teutsch et al., 2005). The apparent isotope fractionation between Fe²⁺(aq) and Fe³⁺(reac) is identical regardless of bacteria species or type of oxides yielding Δ⁵⁶Fe_{Fe(II)-Fe(III)_{reac}} values of ~ -3‰ (Crosby et al., 2007). When the reduction continues, increased Fe²⁺ concentrations in solution result in a shift to a heavier Fe isotopic composition of dissolved Fe because of isotopic pool-size effects. This fractionation depends on the relative proportion of the three main reactive Fe pools (Crosby et al., 2007). Schuth et al. (2015) conducted incubation experiments under controlled E_h conditions and found strong variations in bulk δ⁵⁶Fe values in soil solutions of a surface soil horizon, while a permanently anoxic lower soil horizon showed no Fe release and no response of δ⁵⁶Fe values to flooding, probably because of a more stable crystalline Fe oxide mineralogy or an absence of Fe-reducing bacteria. However, at constant E_h a steady state of δ⁵⁶Fe values and dissolved total Fe was observed by Schuth et al. (2015) for the topsoil because of the balance of Fe release and removal from solution. Garnier at al. (2017) and Thompson et al. (2007) found Rayleigh-type fractionation during DIR in redox-influenced soils but smaller fractionation compared to the experimentally determined fractionation factors, because of additional processes occurring in soil (e.g. organic complexation).
Ilina et al. (2013a) studied iron isotope variations in river, lake and mire waters and soil solutions of different filtration sizes reaching from 100 µm to <1 kDa. The $\delta^{57}\text{Fe}$ values of mire waters (0.12 to 0.66 ‰) and soil solutions (-0.20 to -0.42 ‰) showed no systematic differences except for the <1 kDa fraction, which was the lightest (-0.83 to -0.91 ‰). River waters became constantly heavier (up to 4.2 ‰) with decreasing pore size, which was explained by stronger Fe binding of low molecular weight organic compounds compared to bigger particles like Fe (oxyhydr)oxides (Ilina et al., 2013a). A similar observation was made by Schuth and Mansfeldt (2016) for stagnant water, where smaller particles indicated a slight enrichment in $^{56}\text{Fe}$. Furthermore, other mechanisms can induce Fe isotope fractionation. Heavy Fe isotopes are preferentially adsorbed or precipitated on the bacterial cell surface ($\Delta^{56}\text{Fe}_{\text{cell-solution}}$ up to 1.7‰) and complexed by organic ligands ($\Delta^{56}\text{Fe}_{\text{complex-solution}}$ of 0.6‰; Brantley et al., 2001; Dideriksen et al., 2008; Morgan et al., 2010; Mulholland et al., 2015). Precipitation of siderite caused a lighter Fe isotope ratio in the siderite ($\Delta^{56}\text{Fe}_{\text{dissolved-siderite}} = 0.0-0.9‰$; Johnson et al., 2005; Wiesli et al., 2004). While kinetic FeS precipitation caused an enrichment of light Fe in the precipitate ($\Delta^{56}\text{Fe}_{\text{Fe(II)aq-FeS}} = 0.85\pm0.30‰$; Butler et al., 2005), precipitation of mackinawite at equilibrium conditions caused oppositional fractionation of $\Delta^{56}\text{Fe}_{\text{Fe(II)aq-mackinawite}} = -0.32\pm0.29‰$ to -0.64±0.36‰ (Guilbaud et al., 2011; Wu et al., 2011). The precipitation of magnetite similarly causes an enrichment of heavy isotopes in the precipitates ($\Delta^{56}\text{Fe}_{\text{Fe(II)aq-magnetite}} = -1.34\pm0.11‰$ to -1.56±0.20‰; Friedrich et al., 2014; Johnson et al., 2005). Sorption of Cu to Al and Fe (oxyhydr)oxides causes an enrichment of heavy Cu on the surface of these minerals with $\Delta^{65}\text{Cu}_{\text{solid-solution}}$ values of up to 1‰ (Balistrieri et al., 2008; Pokrovsky et al., 2008). Sorption to organic ligands shows varying fractionation with $\Delta^{65}\text{Cu}_{\text{solid-solution}}$ values between -0.3 and 0.8‰ depending on type of organic ligand and pH (Bigalke et al., 2010a; Navarette et al., 2011; Ryan et al., 2014). Lighter Cu isotopes are preferentially adsorbed on clay mineral surfaces with $\Delta^{65}\text{Cu}_{\text{adsorbed-solution}}$ values of -0.3‰ (Li...
Copper uptake into bacteria also causes various Cu isotope fractionations depending on organism and pH. Mainly the lighter Cu isotope is preferentially incorporated into the live bacteria cell ($\Delta^{65}\text{Cu}_{\text{bacteria-solution}} = 0$ to $-4.4\%$; Navarette et al., 2011; Pokrovsky et al., 2008; Zhu et al., 2002). In redox reactions, the reduced Cu species is enriched in lighter Cu isotopes (e.g., $\Delta^{65}\text{Cu}_{\text{solution-covellite}} = 3.06\%$; Ehrlich et al., 2004 and $\Delta^{65}\text{Cu}_{\text{Cu}^{2+}-\text{CuI}} = 4\%$; Zhu et al., 2002). Bigalke et al. (2010b, 2011, 2013) found episodically water-saturated soil to be enriched in heavy Cu isotopes, which they attributed to the loss of light Cu by leaching of reduced colloidal Cu forms. Ilina et al. (2013b) found no significant variations ($\delta^{65}\text{Cu}$ values of 0.36 to 0.46 $\%$) between differently filtered fractions (100$\mu$m - <1kDa) of river water, which they attributed to similar bonding strength of Cu fulvic complexes of different mass. Vance et al. (2008) reported lighter $\delta^{65}\text{Cu}$ values (-0.24 to -1.02$\%$) in the particulate fraction of rivers compared to the dissolved phase (0.42 to 0.94$\%$) and explained this with strong bonding of dissolved Cu to organic complexes.

In this study, we investigate the isotope geochemistry of dissolved and colloidal Fe and Cu in a water-saturated carbonatic soil. Our goals are to better understand how a) Fe (oxyhydr)oxide dissolution and Fe colloid formation and b) Cu release and colloid formation in soils affect isotopic composition of soils and soil solutions. We furthermore will link our experimental results to field observations previously reported in the literature.

### 2. Materials and methods

A Calcaric Fluvisol (IUSS, 2014), with high calcium carbonate content (420 g kg$^{-1}$) was sampled near the city of Aesch in Switzerland (47°28'39" N / 7°36'21" E). The soil had a neutral pH (7.4), an organic carbon concentration of 21.7 g kg$^{-1}$, a total Fe concentration of 12.6 g kg$^{-1}$ and a dithionite-extractable Fe concentration of 7.87 g kg$^{-1}$ (Abgottspon et al., 2015). More details about the soil are given in Table S1. The soil was located next to the river Birs on a floodplain with temporary flooding (Kayser et al., 2006). The sampling site was moderately contaminated with several trace metals (e.g.,
Cd, Cr, Cu, Pb, Table S1) by an adjacent non-ferrous metal smelter, causing elevated Cu concentrations of 122 mg kg\(^{-1}\) (Kusonwiriyawong et al., 2016). About 10 kg of topsoil were sampled from the Ah horizon (0-0.15 m depth) in March 2011. The soil was air-dried and sieved to <2 mm. The setup of the incubation experiment was similar to that of Weber et al. (2009b) and identical to that described in Abgottspon et al. (2015). In fact, samples were taken from exactly the same experiment described in Abgottspon et al. (2015). In summary, 1.7 L of artificial river water were filled in 3-L PE microcosms. The artificial river water (prepared from ultrapure water and clean salts) had a similar ionic strength and ionic composition as the river Birs next to the sampling site (0.92 mmol L\(^{-1}\) CaCl\(_2\), 0.98 mmol L\(^{-1}\) NaNO\(_3\), 0.84 mmol L\(^{-1}\) MgSO\(_4\)), but did not contain DOC, trace metals, or suspended matter. Then, 1.7 kg of air-dry soil was slowly added to the microcosm and stirred to ensure homogenization and removal of air bubbles, resulting in complete water saturation. After homogenization, the soil material was allowed to settle and around 2 cm of stagnant water covered the soil (Figure S1). Three replicate microcosms and one blank (i.e. only artificial river water) were set up in a glovebox (GS Glovebox Plexiglas, GS Glovebox Systemtechnik, Malsch, Germany) that was purged with nitrogen (Figure S2). The soil solution was extracted with PE suction cups with a 10-μm nominal pore size (ecoTech GmbH, Bonn, Germany) manually located in the lower half of the microcosms with a syringe. On each sampling date, 50 mL of solution was extracted. Over the whole experiment, we extracted 700 mL of a total solution volume of 1700 mL. Nevertheless, in the lower part of the microcosm the soil was always water-saturated (and the suction cup at this depth always flooded). Our extraction of solution might have simulated slow drainage as is also observed in natural soils. We realized mass balance calculations to check how repeated sampling of the microcosms may affect Fe and Cu concentrations and δ\(^{56}\)Fe and δ\(^{65}\)Cu values. Over the whole experiment we extracted 0.06% and 0.18% of the total Fe and Cu in the system, respectively. Taking the isotopic composition into account, the isotopic change of the total pool related to the sampling is insignificant (-0.001‰ and 0.002‰) for δ\(^{56}\)Fe and δ\(^{65}\)Cu, respectively. These effects are much smaller than our analytical precision. Therefore, we conclude that the sampling will not significantly affect the isotopic composition of the total metal pools. The Eh and pH values were analysed directly after sampling in the glovebox. To rule out that the initial mobilization of colloids is an artifact caused by manually
sampling of the soil solution, Al concentrations were analyzed and found to be consistently low (on
average 0.5 µmol L\(^{-1}\)). As Al is a major constituent of the soil (2.6 wt. %), low Al concentrations
render the artificial release of soil colloids (e.g., clay minerals, Al hydroxides) unlikely. The solution
is supposed to contain the total (colloidal + dissolved) element concentration (<10 µm). To determine
dissolved element concentrations, samples were manually filtered to <0.02 µm using syringe filters
(Anotop 25 Plus, Whatman, Bottingen, Switzerland) in the glove box directly after sampling. The Fe
and Cu concentrations were analyzed by ICP-MS (7700x, Agilent, Santa Clara, California) using In
and Rh as internal standards. The difference between total and dissolved elemental concentrations was
assumed to be the colloidal fraction (0.02-10 µm). All unfiltered and filtered solutions were acidified
with suprapur HNO\(_3\) (ROTH, Karlsruhe, Germany) and stored at -20°C prior to isotope analysis. The
Fe\(^{2+}\) concentration in the unfiltered solution was determined photometrically (Spectroquant Pharoh 100,
MERCK). Each 5-mL aliquot of unfiltered soil solution was treated with 50 µL of H\(_2\)SO\(_4\) (25%, p.a.),
0.5 mL of ammonium-acetate-acetic acid (400 g L\(^{-1}\) ammonium acetate, 50% acetic acid) and 0.2 mL
of phenanthroin solution (5 g L\(^{-1}\) 1,10-phenanthroliumchlorid) and made up to 10 mL with Millipore
water in the glovebox. Samples were measured after 15 minutes at the wavelength of 510 nm. A
detailed evolution of Eh, pH, SO\(_4^{2-}\), S\(^2-\), Fe\(^{2+}\) C and metal concentrations in 2-4 day resolution is given
in Tables S2 & S3 and Abgottspon et al. (2015).

For total digestion of the bulk soil sample, approximately 70 mg of soil was digested
in HNO\(_3\), HF and H\(_2\)O\(_2\) (3:2:1) in PFA beakers (Savillex, MN, USA) at least 24 h on a hot
plate at 120 °C. The digested samples were evaporated until dryness at 70 °C. Soil solution
samples were evaporated in PFA vials (Savillex, MN, USA). The dried residues were digested
using a mixture of concentrated HNO\(_3\) and H\(_2\)O\(_2\) (ratio 1:1) at 120°C for at least 3-4 hours and
evaporated to dryness. The dried soil and solution samples were subsequently re-fluxed in
HCl and HNO\(_3\) (3:1) at least 3 h at 120 °C and finally dissolved in 7 mol L\(^{-1}\) HCl + 0.001%
H\(_2\)O\(_2\) for ion exchange separation. The total (<10µm) and dissolved (<0.02µm) fractions were
analyzed in samples from Days 0, 2, 4, 6, 8, 11, 13, 16, 19, 22, 26, 30, 34 and 39. After 8 days
of incubation, the dissolved Cu concentrations (<0.02µm) were too low to get the required
300 ng of Cu for the isotope analysis. Thus, samples from the dissolved fraction of days 8, 10, 13, and 16 were combined to a single composite sample and another composite sample was prepared from days 19, 22, 26, 30, 34 and 39. In these composite samples of the dissolved fraction, we also measured stable Fe isotope ratios after purification (see below). In addition, for Fe isotope measurement samples from the three replicates microcosms were combined to a composite sample of days 4 and 6, respectively, for both total (<10µm) and dissolved (<0.02µm) fractions because of too low mass (< 7µg Fe) of Fe for isotopic analysis at the beginning of the experiment. Copper and Fe fractions were purified as described in Bigalke et al. (2013). For solution samples, a single-step ion-exchange approach was sufficient to separate Cu and Fe from other matrix elements, while bulk soil samples were purified twice for Cu isotope analysis. All Fe and Cu fractions yielded an average recovery of 100.6±1.3% (mean±SD) and 101.1±2.1% (mean±SD), respectively. All samples which did not yield a recovery of within 100 ± 6% were discarded, and the purification process was repeated. The acids (HNO₃ and HCl) used in this study were purified by sub-boiling distillation in quartz stills or were purchased in suprapur quality. Other reagents were of suprapur quality. All solutions and dilute acids were prepared using 18 MΩ grade water (EMD Millipore, MA, USA). All sample preparations were performed in the clean chemistry laboratory at the Institute of Geology, University of Bern. The procedural blank was 3.1±1.8 ng (n=3) for Fe and 2.3±0.2 ng (n=3) for Cu, compared to at least 7000 and 300 ng of Fe and Cu, respectively, in the samples.

Iron and Cu isotope measurements were performed on a Thermo-Scientific Neptune Plus MC-ICP-MS at the Leibniz University Hannover, Germany. The Fe isotopic compositions were analyzed in the high mass resolution mode to resolve isobaric interferences of \(^{40}\text{Ar}^{14}\text{N}^+\), \(^{40}\text{Ar}^{16}\text{O}^+\), and \(^{40}\text{Ar}^{16}\text{OH}^+\) on \(^{54}\text{Fe}^\text{6+}\), \(^{56}\text{Fe}\), and \(^{57}\text{Fe}\) (Weyer and Schwieters, 2003). Chromium (\(^{53}\text{Cr}\)) was monitored to correct isobaric interferences of \(^{54}\text{Cr}\) on \(^{54}\text{Fe}\). The standard-sample bracketing method was applied in combination with external
element doping for mass bias correction. For the latter, the 5 mg L\(^{-1}\) Ni standard NIST 986 (National Institute of Standards and Technology, Gaithersburg, MA, USA) was doped to all Fe standards (IRMM-014, Institute for Reference Materials and Measurements, Geel, Belgium) and samples for mass-bias correction (Oeser et al., 2014). The Fe isotope standard (IRMM-014) and the samples were diluted to yield a concentration of 7 mg L\(^{-1}\) Fe with 2% HNO\(_3\). Every sample was at least analyzed twice and the mean \(\delta^{56}\)Fe value is given. The \(\delta^{56}\)Fe values are reported relative to IRMM-014.

In a three-isotope plot, \(\delta^{56}\)Fe and \(\delta^{57}\)Fe values define a linear correlation (slope=1.4664, \(R^2= 0.99\), Figure S3) following the theoretical mass-dependent fractionation line (\(\delta^{57}\)Fe \(\approx\)1.5 x \(\delta^{56}\)Fe) which indicates the absence of isobaric interferences (Malinovsky et al., 2003). The accuracy and precision of the Fe isotope analysis was monitored using an Fe-free matrix sample (BCR-2, Basalt Columbia River 2, USGS, Reston, VA, USA) that was doped with Fe of known isotopic composition, and an in-house standard (an Fe salt from ETH Zurich, Switzerland). The Fe-free matrix samples were prepared from the matrix fraction derived from the purification of the original samples which were doped with the Fe standard (Merck Certipur\(^{®}\), Darmstadt, Germany) and processed in the same way as the samples. The average \(\delta^{56}\)Fe value of the matrix fraction was 0.19±0.03‰ (mean ±2SD, n=5) which is identical to the pure Merck Fe value (0.19±0.02‰, n=3). The USGS basalt BCR-2 yielded a mean \(\delta^{56}\)Fe value of 0.07±0.02‰ (2SD, n=4) which agrees well with previously published values ranging from 0.05±0.02‰ to 0.11±0.03‰ (Craddock and Dauphas, 2011; Liu et al., 2014a; Schuth et al., 2015; Weyer et al., 2005). The average \(\delta^{56}\)Fe value of our in-house standard was -0.69±0.02‰ (mean ± 2SD, n=6) and again in good agreement with previously published values varying from -0.71±0.18‰ to -0.73±0.10‰ (Fehr et al., 2008; Kiczka et al., 2011; Mansfeldt et al., 2012; Schuth et al., 2015).

For Cu isotope analysis, standard-sample bracketing was conducted in combination with external element doping for mass-bias correction. Again the Ni standard NIST 986 was
used for the mass-bias correction by analyzing $^{62}$Ni/$^{60}$Ni (for details see Lazarov and Horn, 2015). The samples were diluted to yield a final concentration of 300 µg Cu L$^{-1}$ and 1000 µg Ni L$^{-1}$ and were measured with MC-ICP-MS in low resolution mode. Every sample was at least analyzed twice and the mean was reported. The $\delta^{65}$Cu values are reported relative to the Cu standard NIST SRM 976 after external mass-bias correction using the exponential law.

The success of the Cu purification and the accuracy and precision of the Cu isotope analysis was verified by using Cu-free matrix samples that were subsequently doped with Cu of known isotopic composition, and a basalt reference material (BCR-2). The Cu-free matrix samples were prepared from the matrix fraction derived from the purification of the original samples and doped with the ERM$^\text{®}$-AE633 Cu isotope standards (Institute for Reference Materials and Measurements, Geel, Belgium), which is isotopically indistinguishable from NIST 976 (Moeller et al., 2012). The doped matrices were treated and purified in the same manner as the original samples. The average $\delta^{65}$Cu value of the matrix samples was $-0.02\pm0.01\%$ (mean ± 2SD, n=3) which is identical to the value of ERM$^\text{®}$-AE633 ($-0.01\pm0.05\%$; Moeller et al., 2012). The BCR-2 reference material yielded a mean of $\delta^{65}$Cu$_{\text{NIST976}} = 0.18\pm0.01\%$ (2SD, n=3), which is in good agreement with previously published data ranging from $0.14\pm0.05\%$ to $0.22\pm0.06\%$ (Moeller et al., 2012). The external reproducibility was evaluated with the in-house Cu standard (NBS C 125-2) yielding an average $\delta^{65}$Cu value of $0.37\pm0.06\%$ (mean ±2SD, n=10).

Most isotopic compositions from our experiments are reported as means of the triplicate microcosm incubations (Table 1). Therefore, error bars in the figures do not reflect measurement uncertainties but variations among the triplicate microcosms reflecting the natural soil heterogeneity. The isotopic composition of the metal colloidal fraction ($\delta_{\text{col}} = \delta^{56}\text{Fe} \text{ and } \delta^{65}\text{Cu values}$) was calculated from the difference in the isotopic composition of total and dissolved metal concentrations in solution based on isotopic mass balance, as shown in Equation 1. To calculate colloidal concentrations and $\delta$ values of the composite samples,
we averaged the values of the colloidal concentrations and the δ values based on mass balance
approaches, as shown in Equation 2 with δ_{ss} and C_{ss} being the delta value and the
concentration of the respective subsample. In Equation 1-5 δ_{tot}, δ_{diss} and δ_{coll} are the δ^{56}Fe or
δ^{65}Cu and c_{tot}, c_{diss} and c_{coll} the concentrations of the total, dissolved and colloidal fraction,
while the prefix 2SD indicate the respective 2SD uncertainty.

\[ \delta_{\text{coll}} = \frac{(\delta_{\text{tot}} \times [c_{\text{tot}}]) - (\delta_{\text{diss}} \times [c_{\text{diss}}])}{[c_{\text{tot}}] - [c_{\text{diss}}]} \]  
\[ \delta_{\text{bulk}} = \frac{\sum_{i=1}^{n} \delta_{\text{ss}} \times [c_{\text{ss}}]}{\sum_{i=1}^{n} [c_{\text{ss}}]} \]  

The error of the δ value of the colloidal fraction calculated according to the general formula
of error propagation (Taylor, 1997) as shown in Equation 3.

\[ 2SD\delta_{\text{coll}} = \sqrt{\left(\frac{\partial \delta_{\text{coll}}}{\partial \delta_{\text{tot}}} 2SD\delta_{\text{tot}}\right)^2 + \left(\frac{\partial \delta_{\text{coll}}}{\partial c_{\text{tot}}} 2SDc_{\text{tot}}\right)^2 + \left(\frac{\partial \delta_{\text{coll}}}{\partial \delta_{\text{diss}}} 2SD\delta_{\text{diss}}\right)^2 + \left(\frac{\partial \delta_{\text{coll}}}{\partial c_{\text{diss}}} 2SDc_{\text{diss}}\right)^2} \]  

If we apply Equation 3 to Equation 1, we get Equation 4, which allows calculating the
uncertainty of the δ values of the colloidal fraction as displayed in Table 1.

\[ 2SD\delta_{\text{coll}} = \sqrt{\left(\frac{c_{\text{tot}}}{c_{\text{tot}} - c_{\text{diss}}} 2SD\delta_{\text{tot}}\right)^2 + \left(\frac{c_{\text{diss}}(\delta_{\text{tot}} - \delta_{\text{diss}})}{(c_{\text{tot}} - c_{\text{diss}})^2} 2SDc_{\text{tot}}\right)^2 + \left(\frac{c_{\text{diss}}}{c_{\text{diss}} - c_{\text{tot}}} 2SD\delta_{\text{diss}}\right)^2 + \left(\frac{\delta_{\text{tot}} - \delta_{\text{diss}}}{c_{\text{diss}} - c_{\text{tot}}} c_{\text{tot}}^2 2SD\delta_{\text{diss}}\right)^2} \]  

Finally, the error of the Δ_{diss-coll} Values was calculated according to equation 5

\[ 2SD\Delta_{\text{diss-coll}} = \sqrt{(2SD\delta_{\text{diss}})^2 + (2SD\delta_{\text{coll}})^2} \]  

Table 1 Concentration and isotopic compositions of Fe and Cu in bulk soil prior to flooding, total (<10µm), dissolved (<0.02 µm) and (calculated) colloidal (0.02-10 µm) fractions of soil solution during 40 days of experimental flooding.

<table>
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<tr>
<th>Element</th>
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<th>Dissolved</th>
<th>Colloidal</th>
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<td>δ value (‰)</td>
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⁹Number of replicate preparations and analysis, bConcentrations in the solid soil are in µmol kg⁻¹, cBulked sample combined from days 8 to 16, dBulked sample combined from days 19 to 39, eCalculated by the difference between the δ⁶⁵Cu values of total and dissolved Cu (Eq. 1), fCalculated by error propagation according to Eq. S1-S4, gCalculated by substracting the dissolved from the total concentrations.
3. Results

Iron was first mainly released in colloidal form, which changed after 19 days to predominantly dissolved Fe (Figure 1 and Table S3, Abgottspon et al., 2015). On day 39, only 22% of the total Fe was still in colloidal form. The sum of total and dissolved Fe concentrations increased continuously until the end of the incubation, while the colloidal concentrations increased until day 16 and decreased afterwards (Figure 1). From day 2 on, the total Fe in solution was dominated by Fe²⁺, but the concentration of Fe²⁺ started to deviate substantially from that of total Fe from day 13 on. After day 13, the Fe²⁺ concentrations showed only a slight increase until day 22 and a decrease afterwards. From day 22 on, the dissolved Fe concentration was even higher than that of total Fe²⁺ indicating that Fe³⁺ was also present in the dissolved phase. On day 29, only 33% of the total Fe in solution was still Fe²⁺.

The Fe isotope composition of total Fe in solution started at negative δ⁵⁶Fe values on day 4 and afterwards increased slightly (Figure 2b). The δ⁵⁶Fe value of total Fe in solution was correlated with the total Fe concentrations in solution (Figure 3). The δ⁵⁶Fe value of the colloidal Fe sharply increased on day 6 and then gradually continued to increase until the end of incubation, and was parallel to the evolution of the δ⁵⁶Fe values of the total Fe in solution (Figure 2b). The δ⁵⁶Fe values of the dissolved fraction were higher than the colloidal fraction on day 4. Then, δ⁵⁶Fe values of the dissolved fraction continuously decreased from day 4 to day 8-16. After 19-39 days, the δ⁵⁶Fe values of dissolved Fe shifted to heavier values (Figure 2b). The Δ⁵⁶Fe_{dissolved-colloidal} value decreased from 0.31±0.04‰ on day 4 to -1.86±0.26‰ on day 12 and thereafter increased slightly (Figure 4a). The δ⁵⁶Fe value of the bulk soil was 0.04±0.01‰ (n=3).

Copper was also mainly released in colloidal form but in contrast to Fe, Cu release peaked around day 6 and decreased thereafter (Figure 2c, Abgottspon et al., 2015). The δ⁶⁵Cu values of the total, dissolved and colloidal Cu fractions did not differ from each other and
were close to that of the bulk soil at the start of the experiment (day 0; Figure 2d). With strongly increasing Cu concentrations in solution on day 2, the $\delta^{65}\text{Cu}$ value of total Cu in solution shifted to lower values and later on back towards initial values (Figure 2d). The $\delta^{65}\text{Cu}$ value in both colloidal and dissolved Cu decreased markedly on day 2. After day 2, the $\delta^{65}\text{Cu}$ value of the colloidal Cu increased until day 16; thereafter the $\delta^{65}\text{Cu}$ value decreased slightly until the end of the experiment. The $\delta^{65}\text{Cu}$ value of the dissolved fraction increased as well until day 6, but then, decreased towards the end of the experiment. The $\Delta^{65}\text{Cu}_{\text{dissolved}-\text{colloidal}}$ value changed from a maximum of $1.58\pm0.13\%$ on day 6 back to $0.81\pm0.03\%$ at the end of the experiment (Figure 4b). The $\Delta^{65}\text{Cu}_{\text{dissolved}-\text{colloidal}}$ from day 0 of the experiment could not be used because of the high uncertainty of the $\delta^{65}\text{Cu}_{\text{coll}}$ value at this day (Tab. 1). The $\delta^{65}\text{Cu}$ value of the bulk soil was $-0.08\pm0.04\%$ (n=3).

Figure 1 Fe concentrations in total, dissolved and colloidal Fe during the experiment. Total, dissolved and colloidal Fe concentrations were taken from Abgottspon et al. (2015). The Fe$^{2+}$ concentrations were specifically measured for this study in unfiltered samples and therefore represent total Fe$^{2+}$ in solution including dissolved an colloidal fractions.
Figure 2 Temporal variation in a) Fe concentration, b) Fe isotope compositions, c) Cu concentration, and d) Cu isotope compositions during 40 days of flooding. Connection lines of the single data points are displayed to guide the eye. Plateau-like connection lines for Fe and Cu are displayed because of the pooling of the samples from different days. So every step corresponds to one sample. In Panels b and d, the dashed horizontal lines show the δ^{56}Fe (0.04±0.01‰) and δ^{65}Cu (-0.08±0.04‰) values of the bulk soil, respectively. Vertical bars in Panels a and c indicate SD of concentrations from triplicate microcosm experiments. Vertical bars in Panels b and d indicate 2SD for isotope values. Error bars are smaller than the symbols unless visible. Please note the different scales.
4. Discussion

4.1. Fe isotopes

After the flooding of soils, oxygen is consumed and electrons are released by microorganisms gaining energy from oxidation of organic matter. When the dissolved oxygen is used up, the electrons are transferred to alternative electron acceptors (Lovley, 1991). Iron (oxyhydr)oxides become the dominant alternative electron acceptors in flooded soils at sufficiently low $E_h (< 150 \text{ mV})$. In our experiment, the $E_h$ value decreased quickly, reached 90 mV four days after flooding and stabilized around 0 mV afterwards (Abgottspon et al., 2015). This decrease potentially triggers DIR, which is reflected in the temporal course of the total Fe concentration and $\delta^{56}\text{Fe}$ value in solution in our experiment (Figure 3) and agrees with findings of Crosby et al. (2007), Johnson et al. (2005), but also with abiotic reduction reactions (Wiederhold et al., 2006).
Figure 3 $\delta^{56}$Fe values of total Fe in solution as a function of Fe concentrations during 40 days of flooding. Bars indicate SD of Fe concentrations from triplicate microcosm experiments and 2SD of Fe isotope compositions. Error bars are smaller than the symbols unless visible. The dotted line represents a logarithmic trend line.

From days 2-10, nearly all Fe in solution was Fe$^{2+}$, while the contribution of Fe$^{2+}$ to total Fe in solution decreased after day 10 (Fig. 1), which is similar to the increase in Fe$^{3+}$ concentrations in solution reported by Schuth et al. (2015). After day 19, total dissolved Fe showed higher concentrations than Fe$^{2+}$, indicating the presence of Fe$^{3+}$ even in the dissolved phase. Because the solubility of Fe$^{3+}$ is very low, this finding may only be explained by small Fe$^{3+}$ colloids, which passed through the 0.02 µm filter (and are therefore operationally defined as dissolved species), or by the complexation of Fe$^{3+}$ by organic substances, forming soluble complexes. Another possible explanation might be an analytical complication in the determination of the Fe$^{2+}$ concentrations. If the Fe colloids at the end of the experiment did not fully dissolve when the sulphuric acid was added, the Fe$^{2+}$ concentration might have been underestimated.

The pooling of the samples from different days, which was necessary because of the low Fe concentrations at the beginning of the experiment and low dissolved Cu concentrations at the end of the experiment, limit the temporal information of the Fe data. Therefore, we focus the further discussion on the Fe colloid release under anoxic conditions in soils, a process for which no Fe isotope data have yet been reported. The colloidal release plays an important role in our carbonatic floodplain soil. Up to 90% of the Fe was released in colloidal form on day 6 (Table S3; Abgottspon et al., 2015). Even in the later phase of the experiment (after day 10), colloidal Fe still accounted for 20-50% of total Fe in solution (Figure 2a). The lack of colloidal organic carbon and Al in solution (Abgottspon et al., 2015) indicate no co-mobilisation with OM or colloidal soil constituents like clay minerals, but rather Fe to be the main component of the
colloids (Table S2, S3). At the beginning of the experiment, total Fe and Fe$^{2+}$ concentrations in solution were identical, highlighting that colloids consisted of reduced Fe and are not Fe (oxyhydr)oxides released into soil solution (Figure 1). The colloidal Fe pool had low $\delta^{56}$Fe values at the beginning, hence supporting the idea of precipitation from reduced light Fe$^{2+}_{aq}$ rather than mobilization by deflocculation of ferrihydrite, which should have a more positive value (span of the amorphous Fe oxide fraction in hydromorphic and oxic soils approx. -0.7 to 0.4 ‰; Guelke et al., 2010; Schuth et al., 2015; Wiederhold et al., 2007a; Wiederhold et al., 2007b) compared to the colloidal fraction on day 4 of our experiment (-1.16±0.04‰). Indeed, the initial $\Delta^{56}$Fe$^{\text{dissolved-colloidal}}$ value was positive and fitted the values reported for siderite precipitation in laboratory experiments ($\Delta^{56}$Fe$^{\text{dissolved-siderite}}$ values between 0.0 and 0.9‰; Johnson et al., 2005; Wiesli et al., 2004) and similar to that of lake sediments with supposed siderite precipitation (Figure 4a, Teutsch et al., 2009). However, when looking at the proposed $\Delta^{56}$Fe$^{\text{dissolved-colloidal}}$ value, it has to be kept in mind that isotopic fractionation of Fe in solution can also be influenced by interactions with the solid Fe pool in soil. Dissolved Fe$^{2+}$ released by DIR might be sorbed to Fe (oxyhydr)oxide surfaces in soil and undergo electron transfer-atom exchange reactions, a process which influences the $\delta^{56}$Fe value in solution (Crosby et al., 2005; 2007) and might also affect the $\Delta^{56}$Fe$^{\text{dissolved-colloidal}}$ values.
Figure 4 Temporal variations in apparent isotopic fractionation between dissolved and colloidal a) Fe and b) Cu. Horizontal error bars represent the time interval of the pooled samples. Vertical error bars represent the 2SD calculated according to Equations 3-5.
Abgottspon et al. (2015), suggested siderite precipitation from day 8 on in the same experiment. However, there is a time gap between the positive $\Delta^{56}\text{Fe}_{\text{dissolved-colloidal}}$ on day 4 and the proposed siderite precipitation. Furthermore, a confirmation of siderite formation or the precipitation of other minerals by XRD analysis was not possible because of the low mass of colloidal material on the filters and interference of the filter material (an alumina-based membrane).

Siderite can be formed by the reaction between $\text{Fe}^{2+}$ produced by DIR and $\text{HCO}_3^-$ produced by microbial oxidation of organic matter (Mansfeldt et al., 2012; Wiesli et al., 2004). In our soil, high concentrations of inorganic carbon in solution (10 mmol L$^{-1}$ on day 4, Abgottspon et al., 2015) might cause a precipitation of $\text{FeCO}_3$ directly from dissolved $\text{Fe}^{2+}$ and $\text{HCO}_3^-$ in solution at the beginning of the experiment. This is in agreement with the presence of colloidal inorganic carbon between days 2-19 of the experiment (Abgottspon et al., 2015), a dominance of colloidal $\text{Fe}$ and the fact that total $\text{Fe}$ in solution was $\text{Fe}^{2+}$ until day 10 of the experiment. Therefore, even if we have no spectroscopic proof for siderite formation, we consider siderite to be the most likely $\text{Fe}$ mineral in colloidal $\text{Fe}$ based on solution chemistry, isotope results and previous findings (Abgottspon et al., 2015; Du Laing et al., 2009; Mansfeldt et al., 2012; Wiesli et al., 2004). However, already on day 6, the $\Delta^{56}\text{Fe}_{\text{dissolved-colloidal}}$ changed, which may be explained by the following scenarios:

1. **Changing colloid mineralogy.** Mineralogy of the colloids may change with time, depending on changing solution chemistry and microbial community (Fulda et al., 2013; Roh et al., 2003). As an example, siderite and magnetite are formed in similar environments depending on the composition of the surrounding air and dissolved gases, the presence or absence of the carbonic acid-hydrogencarbonate buffer, pH, incubation temperature, type of
substrate, incubation time, and bacteria species (Roh et al., 2003). Sulfate reduction might induce formation of Fe sulfides (Borch et al., 2010). The change from Fe$^{2+}$ to Fe$^{3+}$ in the soil solution and thus changing proportions of Fe$^{2+}$ and Fe$^{3+}$ in the dissolved and colloidal phases might explain the variations in the $\Delta^{56}$Fe$_{\text{dissolved-colloidal}}$ values. Furthermore, the big gap between total Fe and total Fe$^{2+}$ indicate that colloids change from a Fe$^{2+}$ dominated mineral to a mineral containing also Fe$^{3+}$. Observed $\Delta^{56}$Fe$_{\text{dissolved-colloidal}}$ values are consistent with those expected for a number of Fe minerals like e.g., mackinawite or magnetite (Friedich et al., 2014; Guilbaud et al., 2011; Johnson et al., 2005; Wu et al., 2011).

2. Sorption and electron transfer-atom exchange reactions. The shift to lower $\Delta^{56}$Fe$_{\text{dissolved-colloidal}}$ values can also be caused by sorption and possibly electron transfer-atom exchange reactions at the surface of the colloids or soil components (Crosby et al., 2005; 2007, Liu et al., 2015). The divalent dissolved Fe can be sorbed on these surfaces and electron transfer-atom exchange reactions may occur, which change isotopic signatures of Fe in solution (e.g., Friedich et al., 2014). Sorption and electron transfer-atom exchange reactions on colloid surfaces would increase $\delta^{56}$Fe values in colloids and decrease $\delta^{56}$Fe values in solution, which is consistent with our results until day 16. However, also sorption to Fe oxy(hydr)oxide surfaces in the soil and electron transfer-atom exchange reactions, might explain a significant part of the observed $\delta^{56}$Fe variation in the solution (Reddy et al., 2015). After 16 days of flooding, the shift of dissolved Fe to higher $\delta^{56}$Fe values can be attributed to an isotopic pool size effect caused by proceeding DIR and the growth of the dissolved relative to the colloidal Fe fraction (Figure 2b).
However, the changing $\delta^{56}\text{Fe}$ values together with the increasing or constant colloid concentration, indicated that new Fe colloids were formed throughout, while the concentrations of Cu colloids steadily decreased probably because of aggregation and sedimentation.

In a recent experiment, the same soil was incubated under the same experimental conditions and changes in the partitioning of Fe into five operatively defined fractions (F1–F5; NH$_4$NO$_3$-extractable, NaOAc-extractable, NH$_4$Ox-extractable, hot H$_2$O$_2$/NH$_4$OAc-extractable and residual fractions, respectively) were investigated (Abgottspon et al., 2015). At the beginning of the experiment 81% of the Fe in the soil was in the residual fraction which is supposed to consist mainly of silicate-bound Fe and crystalline Fe (oxyhydr)oxides, the second important fraction was the oxalate-extractable fraction (15%, amorphous Fe oxides), while the other fractions had minor importance (< 3%). Over the time of the incubation, Fe concentrations in F1-F4 increased, indicating a substantial redistribution of Fe, which is only partly mirrored in the solution chemistry.

A variation in solid phase $\delta^{56}\text{Fe}$ values of up to 0.93‰ was reported for soils which are seasonally or permanently water-saturated (Fekiacova et al., 2013; Liu et al., 2014b; Mansfeldt et al., 2012; Schuth et al., 2015; Wiederhold et al., 2007a). This can be attributed to the mobilization of reduced isotopically light Fe, which might be leached, leaving the soil enriched in heavier Fe isotopes (Schuth and Mansfeldt, 2016). The fractionation follows an apparent Rayleigh-type behavior with smaller fractionation factors in soils compared to experimentally found values, because of additional processes (e.g., organic complexation and reprecipitation) occurring in the complex soil system (Garnier et al. 2017, Thompson et al., 2007). In contrast, most soils that developed under oxic conditions only display a small variation in bulk solid phase $\delta^{56}\text{Fe}$ values of 0.08‰ (Podzols show variations in $\delta^{56}\text{Fe}$ values of up to 0.6‰; Poitrasson et al.,
In agreement with these findings, we observed a release of light Fe into the soil solution ranging from -1.1‰ to -0.4‰ (Figure 2b). The evolution of the δ⁵⁶Fe values in total Fe during our experiment showed a time dependency with low δ⁵⁶Fe values shortly after flooding which successively approached bulk soil values with increasing time of flooding.

These findings agree with the interpretation of the variation in δ⁵⁶Fe values of total Fe in the water of the Rio Negro River, where a substantial temporal variation was observed (dos Santos Pinheiro et al., 2014). In Rio Negro River water, the lowest δ⁵⁶Fe values were coupled to strong rain events, which cause a reductive mobilization of Fe from the soils in the river catchment (Bergquist and Boyle, 2006). Hence, colloidal Fe release in certain anoxic soils significantly affects the temporal isotopic variability of the colloidal fraction. For a detailed understanding of the processes causing the isotopic changes in the colloid fraction, further investigation of the associated processes and mineralogy will be needed which is beyond the scope of this study.

Overall, the agreement of the results from our microcosm experiment and field observations confirms that such experiments can be used for future research into Fe isotopic behavior in anoxic soils. A big advantage of the microcosms is that soil solution samples can be extracted under controlled oxygen-free conditions and allow e.g., for the separation of colloids from dissolved Fe in anoxic samples or photometric Fe²⁺ analysis. This separation is hardly possible under oxic field conditions as reduced Fe²⁺ will almost immediately precipitate as Fe (oxyhydr)oxides when coming in contact with the atmosphere. In spite of the limitations of our study because we had to pool bulk samples of several dates and the lack of direct characterization of the mineral phases, our results indicate that the Fe release in colloidal form
contributes substantially to total Fe concentrations and $\delta^{56}\text{Fe}$ values in solution and thus to Fe mobility in our study soil.

4.2. Cu isotopes

The initial apparent isotopic fractionation between Cu in solution (total, dissolved and colloidal) and in bulk soil was small (Figure 2d). With increasing time after flooding, the fractionation between bulk soil and solution and between dissolved and colloidal Cu species in solution increased, with total and colloidal fraction becoming lighter and the dissolved fraction becoming heavier. Relating the Cu isotope ratios to a similar experiment in which Cu speciation was investigated by XAS on a polluted soil from the River Mulde in Germany, offers a possible explanation for most of the observed isotope fractionations. The Cu concentration peak on day 2 and the strong shift to negative $\delta^{65}\text{Cu}$ values of total Cu in solution could be caused by Cu-carrying bacteria cells which were dominating the colloidal phase in studies with other soils (Hofacker et al., 2013; Weber et al., 2009a). The simultaneously observed heavy Cu isotope ratio of the remaining Cu in solution would then be the consequence of previous preferential uptake of $^{63}\text{Cu}$ by the bacteria (Navarette et al., 2011; Pokrovsky et al., 2008), Cu reduction in the cell and the change to trigonal coordination (Weber et al., 2009a). The shift in $\delta^{65}\text{Cu}$ values between dissolved and colloidal Cu was similar to previously reported Cu isotope fractionation that was linked to the uptake of Cu into bacteria with $\Delta^{65}\text{Cu}_{(\text{solution-bacteria})}$ values between $+1.0\%$ and $+4.4\%$ (Navarette et al., 2011). The change of $\delta^{65}\text{Cu}$ values in solution between days 2 and 6, while total Cu concentrations remained stable, can only be explained by the exchange of solution Cu with solid soil Cu pools. The shift to higher $\delta^{65}\text{Cu}$ values from $-1.01\pm0.04\%$ to $-0.50\pm0.13\%$ in total Cu in solution requires the removal of isotopically light Cu from solution and/or release
of heavy Cu into solution. In the experiments of Hofacker et al. (2013) and Weber et al. (2009a), bacteria released Cu\(^+\), which disproportioned into Cu\(^{2+}\), released into solution, and zero-valent Cu (Cu\(^0\)) colloids, in this phase of the incubation. In solution, Cu\(^{2+}\) released from bacteria was partly reduced and precipitated together with the already present Cu in soil solution and exchangeable Cu from the solid soil as Cu\(_x\)S (Weber et al., 2009b). If this process also occurred in our experiment the Cu\(_x\)S would be a mixture of soil Cu and solution Cu and would probably carry a heavier isotopic signal than that of the Cu in the bacteria. After day 6 with the onset of sulfate reduction (Table S2, Abgottspon et al., 2015), Cu isotope composition and apparent fractionation between dissolved and colloidal Cu stabilized (average values of \(\delta^{65}\text{Cu} = -0.29\pm0.15\%\text{o} \) and \(\Delta^{65}\text{Cu}_{\text{dissolved}-\text{colloidal}} = 0.87\pm0.09\%\text{o} \) despite decreasing Cu concentrations in solution. These findings indicate that the further decrease of colloidal Cu was caused by a progressive aggregation and sedimentation of colloids, which does not cause a pronounced isotopic fractionation (Figure 4b). Even if the results from the soil investigated by Hofacker et al. (2013) and Weber et al. (2009a) cannot be directly transferred to our soil because of the acid soil used by these authors and our slightly alkaline soil, Abgottspon et al. (2015) showed that in our soil decreasing metal mobilization paralleled increasing sulfate reduction. However, the fact that we did not observe colloidal organic carbon at the beginning of the experiment (Table S2) disagrees with the findings of bacterial release causing the increase of colloidal Cu at the beginning of the experiment (Hofacker et al., 2013; Weber et al. 2009a). Assuming a C/Cu (w/w) ratio of about 4000 as described for methanogenic bacteria (Scherer et al., 1983), a bacterial release causing 23 \(\mu\text{mol Cu L}^{-1}\) on day 2 should be clearly visible also in colloidal C (approx. 500 mmol L\(^{-1}\)). Even if the strong variation of Cu concentrations in bacteria is taken into
account, the colloidal C concentration should be clearly higher than the measured 0.2 mmol L\(^{-1}\) \(C_{\text{coll}}\) (Table S2), if bacteria caused the colloidal Cu concentration on day 2.

One might hypothesize that the reported changes of Cu redox states (Weber et al., 2009a) explain much of the observed \(\Delta^{65}\text{Cu}_{\text{dissolved-colloidal}}\) values in soil solution, which are 0.82±0.20‰ on day 2, when Cu is supposed to occur as Cu\(^{+}\) and reach a maximum value of 1.58±0.13‰ on day 6 as a result of the presumed dominance of Cu\(^{0}\). After sulfate reduction on days 6-8 (Table S1), \(\Delta^{65}\text{Cu}_{\text{dissolved-colloidal}}\) values changed back to 0.81±0.03‰, in line with the assumption that Cu\(^{+}\) was the dominant Cu species (Weber et al., 2009a; Figure 4b). However, it is not clear, if the same redox transformations as described by Weber et al. (2009a) occur in the soil used in our experiment. Generally, the isotopic fractionation associated with abiotic oxidative leaching of reduced Cu minerals (chalcoite, chalcopyrite, enargite) is in the range of \(\Delta^{65}\text{Cu}_{\text{aq-mineral}} = 0\) to 2.7‰ in laboratory experiments and field surveys (Fernandez and Borrok, 2009; Kimball et al., 2008; Mathur et al., 2005) and agree well with theoretical estimations stating that Cu(I) complexes preferentially enrich \(^{63}\text{Cu}\) and their precipitation results in the enrichment of \(^{65}\text{Cu}\) in the fluid (Fuji et al., 2013). This pattern supports the interpretation that the observed \(\Delta^{65}\text{Cu}_{\text{dissolved-colloidal}}\) values in our experiment are caused by precipitation of reduced colloidal Cu species. However, fractionation has been reported to be irrelevant or even in the opposite direction in the presence of bacteria (Kimball et al., 2008, Mathur et al., 2005). The latter has been explained with the accumulation of heavy isotopes on the bacteria and preferential precipitation of heavy Cu on bacteria surfaces. The biotic induced isotope fractionation is counteracting the isotope fractionation introduced by abiotic redox processes. Thus in a system with reducing conditions and bacteria the \(\delta^{65}\text{Cu}\) values in solution are determined by the balance of Cu released abiotically and Cu that interacts with the cells and biotic precipitates (Mathur et
al., 2005, Kimball et al., 2008). If we transfer the latter findings to our $\Delta^{65}\text{Cu}_{\text{dissolved-colloidal}}$ values, this might indicate that abiotic reduction rather than bacterial Cu precipitation is governing colloid formation. However, the influence of microorganisms as well as other soil process (e.g. organic complexation or reprecipitation as described by Thompson et al., 2007 for Fe), might decrease the apparent fractionation in our soil, compared to experimentally determined fractionation factors in less complex systems (e.g., Fernandez and Borrok, 2009; Zhu et al., 2002).

In a recent experiment (Kusonwiriyawong et al., 2016), we determined the changes in Cu partitioning and the $\delta^{65}\text{Cu}$ values in five fractions of a sequential extraction (F1–F5; NH$_4$NO$_3$-extractable, NaOAc-extractable, NH$_4$Ox-extractable, hot H$_2$O$_2$/NH$_4$OAc-extractable and residual fractions, respectively) in the same soils that we used in this study. We found that in the dry soil, $\delta^{65}\text{Cu}$ values in F1–F4 followed the estimated bonding strengths of Cu in the respective fractions, indicating equilibrium distribution of Cu at the beginning of the experiment. After flooding, Cu concentrations decreased in F1–F3 and increased in F4–F5. Overall, 73% of the total Cu was redistributed among the five studied fractions. The strongest variations in $\delta^{65}\text{Cu}$ values occurred in F3 (0.09±0.07‰ to 1.43±0.13‰) and F4 (−0.24±0.07‰ to 0.55±0.07‰), while flooding had no or small effects on the $\delta^{65}\text{Cu}$ values of F1, F2 and F5. So even if F1 should be the Cu pool, which interacts most strongly with the solution, it did not mirror the shifts of the dissolved fractions in this experiment. The results from Kusonwiriyawong et al. (2016) suggest a direct transfer of Cu from F3 to F4 because both concentration changes and changes in $\delta^{65}\text{Cu}$ values were balanced between the two fractions. The responses of Cu partitioning and $\delta^{65}\text{Cu}$ values to flooding are in line with the formation of CuxS or other reduced Cu species and the reduction of Cu associated with Fe (oxyhydr)oxides.
One factor which can only be traced with the used isotopic approach is the exchange of Cu in solution with solid soil Cu, which is most pronounced between days 2-6. On day 8, the total $\delta^{65}$Cu value of the soil solution showed good agreement with that of Fraction 4 of the sequential extraction of the solid soil by Kusonwiriyawong et al. (2016) on day 7, which is supposed to represent reduced Cu phases. The agreement in these two $\delta^{65}$Cu values supports the assumption, that at this time colloidal Cu in solution and solid soil Cu are coupled. However, the Cu isotope signatures in Fraction 4 in the solid soil changed to positive values (0.55±0.07‰) on day 35, while dissolved Cu still showed negative $\delta^{65}$Cu values and did not change significantly between days 8 and 39. This suggests that after the initial peak Cu colloid concentration, colloid formation and exchange with the soil stopped, while aggregation in solution decreased the Cu colloid concentration.

Babcsanyi et al. (2014) and Bigalke et al. (2010b, 2011, 2013) investigated different oxic and anoxic soils and wetland systems and found that Cu isotope signatures were isotopically heavier in temporally flooded soils than in soils, which developed in an oxic environment. They concluded that isotopically light Cu was released from flooded wetlands. Furthermore, these authors suggested that leaching of reduced Cu colloids was responsible for the observed apparent fractionation. This assumption is confirmed by our findings, which demonstrate a strong release of light colloidal Cu shortly after flooding. Leaching of this Cu from the soil would cause the described shift to heavier $\delta^{65}$Cu values. The extent of the fractionation in real soils would depend on the progress of leaching. In our study soil, the lowest $\delta^{65}$Cu value for total Cu in solution occurred on day 2. Assuming a similar $\delta^{65}$Cu value (-1.0‰) of the total Cu released into solution and leached over several redox cycles and a leaching of e.g., 10% of the initial Cu in bulk soil, the total Cu in bulk soil should be isotopically shifted from initially 0 to +0.1‰.
This mobilization of Cu has to be considered in the management of polluted sites, by preventing changes of redox conditions. Especially, floodplain soils, which show high metal concentrations in many cases, might directly leach the Cu to the associated river, and thus negatively affect the river ecosystem. Because the colloids are carrying up to 100% of Cu and associated trace metals in soil solution and will strongly affect transport behavior and toxicity (Griffitt et al., 2007; Ju-Nam and Lead, 2008), the extent and stability of colloids formed in carbonatic soils need to be considered in risk assessments.

5. Conclusions

- Total $\delta^{56}\text{Fe}$ values in solution followed the pattern of DIR. The $\Delta^{56}\text{Fe}_{\text{dissolved-colloidal}}$ values indicated that dissolved Fe precipitated directly to form colloids, which accounted for 90% of total Fe at the beginning to 20% of total Fe at the end of the experiment. While siderite precipitation might be one reason for colloidal Fe formation, a change of $\Delta^{56}\text{Fe}_{\text{dissolved-colloidal}}$ values with time indicates either changes in the colloid mineralogy, or sorption and possible electron transfer-atom exchange reactions with colloidal Fe or Fe minerals in the soil solid phase. The results match previous findings from lake, river and soil environments and underline the significance of colloid formation from dissolved Fe in some anoxic soil environments. The strong changes in $\Delta^{56}\text{Fe}_{\text{dissolved-colloidal}}$ values raise new questions about formation and changes of the Fe colloid composition under anoxic conditions.

- The $\Delta^{65}\text{Cu}_{\text{dissolved-colloidal}}$ seemed to be mainly driven by Cu redox state. The $\delta^{65}\text{Cu}$ values indicated a strong exchange of colloidal Cu with solid soil Cu and a decoupling of these two pools with beginning sulfate reduction and decreasing Cu concentrations in solution.
The comparison of the experimental results with previous findings from hydromorphic soils and wetlands indicates that Cu colloid formation might be an important factor driving Cu leaching in these environments. Therefore, Cu colloid formation should be accounted for in risk assessment of e.g., polluted floodplain soils and similar environments.

In this study, the first results of Fe and Cu isotopic composition of colloids forming in anoxic soil environments are presented. Both elements are redox sensitive and probably undergo reduction, but show distinctly different temporal concentration and isotopic patterns. Both elements have a considerable colloidal component in common. Colloidal Fe and Cu might cause colloidal co-mobilization of a line of other potential toxic elements in soils. Furthermore, colloidal properties need to be considered in toxicity and mobility assessments.

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Supporting Material
Supporting material for this manuscript is available, including soil characterization, solution chemistry data and figures illustrating the experimental setup and a three isotope plot for the Fe isotope analysis.
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