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Inferences from the vertical distribution of Fe isotopic compositions on pedogenetic processes in soils

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ABSTRACT

The isotopic compositions of major elements in soils can help understand the mechanisms and processes that control the evolution of soils and the nature and dynamics of the soil constituents. In this study, we investigated the variations of the Fe concentrations and isotopic compositions combined with classical soil parameters, such as granulometry, pH, and C and N concentrations. We selected three soils submitted to different hydrodynamic functioning along a toposequence: a well-drained Cambisol and two hydromorphic soils, an Albeluvisol and a Gleysol. In the Cambisol, the isotopic variations were small indicating little redistribution of Fe which we attributed to centimetric-scale exchanges from the Si-bound to the weakly-bound iron pools and insignificant subsurface Fe export. In contrast, the hydromorphic soils showed an overall variation of 0.37‰ for $\delta^{56}$Fe ($\delta^{56}$Fe (‰) = $[(^{56}\text{Fe}/^{54}\text{Fe})_{\text{sample}}/(^{56}\text{Fe}/^{54}\text{Fe})_{\text{IRMM-014}} - 1] \times 1000$) and an inverse correlation between the Fe isotopic compositions and the oxide-bound Fe concentrations. We suggest that, in the uppermost horizon, the mobilisation of oxide-bound Fe was due to the reducing conditions and predominantly involved the light Fe isotopes. Similarly, within the Bt horizon of the Albeluvisol, the fluctuations of the water table level induced changes in the redox conditions and thus Fe dissolution and transport of isotopically light Fe. The Fe isotopic composition profile in the B/C horizon of the Gleysol is dominated by the signature of the parental material. Overall, the variations of the underground water table combined with topography-driven water flow were suggested to be the main mechanisms of Fe translocation in these hydromorphic soils. Finally, the comparison between Fe isotopes profiles in worldwide soils allow us to show that Fe isotopic variations can help discriminate between various mechanisms and scales of Fe transfer in soils and, accordingly, provide information on the evolution of soils, when used in combination with pedological, geochemical, geographical, and environmental characterisations.
Key Words: iron isotopes, soil toposequence, hydromorphy, world soils

Highlights

- This study investigates the Fe behaviour in three soils along a toposequence.
- Fe transport mechanisms were studied using Fe isotopes and classical soil parameters.
- In the Cambisol, Fe transfer is limited; in-situ Fe transformations were dominant.
- In the hydromorphic soils, the oxide-bound Fe controls the Fe transformations.
- Iron isotope ratio variations in soils help distinguish soil evolution processes.
1. Introduction

Understanding mechanisms and processes that control soil evolution, the nature and dynamics of major soil constituents and their behaviour in response to natural and anthropogenic changes is among the key scientific challenges in soil research. Iron is one of the major elements in soils. It is released during the alteration of rocks and soils, and participates in mineral neoformation, thereby playing a crucial role in soil differentiation. Iron concentration in soil varies because of different processes: physico-chemical redistributions without transport and short-distance or long-range transport of Fe. In addition, the fate of Fe influences the chemical cycles of other important elements in soils, such as nutrients (e.g. P, Zn) and pollutants (e.g. As, Cd, Zn) (e.g. Anderson and Christensen, 1988; Jacobs et al., 1970; Ramos et al., 1993). Finally, Fe is an essential micronutrients for plants (Marschner, 1995) and human nutrition (WHO, 2002).

Stable isotopes analyses have proven to be valuable tools to understand biogeochemical processes in soils (e.g. Fry, 2006). Until the end of the 90’s, these studies were limited to C, N, O, H, and S. However, significant analytical developments over the past years, notably due to the advent of Multi-Collector Inductively-Coupled Plasma Mass-Spectrometers (MC-ICP-MS), now allow high-precision analyses required to measure the small isotopic variations of the so-called “non-traditional” stable isotopes that include Fe (e.g. Albarède and Beard, 2004; Dauphas and Rouxel, 2006). Iron has four stable isotopes, $^{54}$Fe, $^{56}$Fe, $^{57}$Fe and $^{58}$Fe. The $\delta$-notation is commonly used to describe the isotopic fractionation relative to the isotopic reference material IRMM-014: $\delta^X$Fe ($\permil$) = 
\[ \frac{([^{X}Fe/^{54}Fe]_{sample}/[^{X}Fe/^{54}Fe]_{IRMM-014} - 1)\times1000, \text{where } X = 56, 57 \text{ or } 58.\]

In natural environment, $\delta^{56}$Fe ranges between -3.5 $\permil$ and +1.5 $\permil$ (Beard et al., 2003a; Beard and Johnson, 2004; Dauphas and Rouxel, 2006). Igneous rocks represent an
isotopically homogeneous terrestrial baseline ($\delta^{56}\text{Fe} = 0.0 \pm 0.05\%$) (Beard et al., 2003a), whereas isotopic heterogeneities were initially measured mostly in rivers and lakes, rocks and minerals formed at low temperature, hydrogenous ferromanganese precipitates, pyrites and hydrothermal systems (e.g. Dauphas and Rouxel, 2006; Fantle and DePaolo, 2004).

Continental weathering involves processes of mechanical and chemical breakdown of rocks and minerals and subsequent soil development. The parent material, i.e., igneous rocks and clastic sediments, show limited isotopic variations: $\delta^{56}\text{Fe} = 0.0 \pm 0.3\%$ (Beard and Johnson, 2004; Fantle and DePaolo, 2004; Johnson et al., 2003). In contrast, soils display a range of variations of the $\delta^{56}\text{Fe}$ from -0.62 to +0.72 ‰ (Emmanuel et al., 2005; Fantle and DePaolo, 2004; Poitrasson et al., 2008; Thompson et al., 2007; Wiederhold et al., 2007a, 2007b) indicating that pedogenic processes, leading to soil formation and evolution, generate Fe isotopes fractionations with respect to the parent material. Experimental studies have shown that mineral dissolution results in the preferential liberation of light Fe isotopes into the solution while the residual material becomes accordingly heavier than the parent material (e.g. Beard et al., 1999; Brantley et al., 2004; Wiederhold et al., 2006). In contrast, sorption of Fe(II) onto goethite and mineral ferrihydrite neoformation arising from the abiotic oxidation of aqueous Fe(II) into Fe(III) appears to favour the heavy isotopes (Bullen et al., 2001; Icopini et al., 2004). These experimental results have been confirmed by soil studies that show that, in general, the mobile fraction in soils has lighter Fe isotopic composition (Brantley et al., 2004; Fantle and DePaolo, 2004; Thompson et al., 2007; Wiederhold et al., 2007a, 2007b). Therefore, the variations of the Fe isotopic signatures in soils can help to investigate the behaviour of Fe in the near-surface environment during pedogenesis and to discriminate between potential processes at the origin of soil evolution.

In this study, we investigated both vertical and lateral variations of the Fe concentrations and isotopic compositions combined with the study of classical soil
parameters such as granulometry, pH, and C and N concentrations. We used three soil profiles submitted to different hydrodynamic functioning along a toposequence: a well-drained oxic profile and two profiles that are waterlogged during a part of the year. The objectives were to (1) study the expression of the isotopic fractionation of iron in these soils, (2) characterize the mechanisms of Fe transport, (3) compare the results with those obtained in previous studies of the Fe isotopes fractionation in soils and (4) evaluate the potential of Fe isotopes to record information about mechanisms of soil transformations.

2. Material and methods

2.1. Study area, sampling and soil description

The sampling area is located in the Kervidy-Naizin catchment that extends over an area of 4.9 km$^2$ in the centre of Brittany, western France and belongs to the Environment Research Observatory (ERO) AgrHyS (response time in Agro-Hydro Systems). The region is characterised by temperate oceanic climate according to the Köppen climate classification, with a mean annual precipitation of 909 mm and a mean monthly temperature ranging from 5.4°C (January) to 17.4°C (August). The land-use is dominated by corn and wheat farming, temporary pastures for dairy production and indoor pig-stock breeding.

The soils in this catchment developed from the weathering of the sedimentary Brioverian schist unit (older than 530 Ma) and eolian Quaternary deposits that overlay locally the bedrock (Olivié-Lauquet et al., 2001; Thomas and Le Berre, 2009; Van Vliet-Lanoe et al., 1998; Walter and Curmi, 1998). The soils are organised in a toposequence consisting of three soil types: (1) a well-drained, cropped Cambisol (IUSS Working Group WRB, 2006) on the upper part of the slope, characterised by oxic conditions, (2) an Albeluvisol (IUSS Working Group WRB, 2006) at midslope, which represents a transition zone between the well-drained cropland and the poorly drained lowermost part of the landscape and (3) a Gleysol (IUSS...
Working Group WRB, 2006) developed next to a small creek that flows at the bottom of the slope. The Albeluvisol and Gleysol are planted with poplar trees. Both soils are roughly ploughed every two years to avoid weed growth between the trees (Durand et al., 1998). They both undergo seasonal fluctuations of water saturation, with winter-spring corresponding to a period of reducing chemical conditions while summer, or late summer in the case of the Gleysol, is dominated by oxidative chemical conditions (Davranche et al., 2011; Trolard et al., 2002). The water table in these two soils is located close to the topographic surface (Davranche et al., 2011; Olivié-Lauquet et al., 2001; Pauwels et al., 1996). During rainy periods a temporary water table could develop in the Ap-horizon and lateral drainage could occur.

The sampled profiles are located close to the sites C, I and F described by Trolard et al. (2002) for the Cambisol, the Albeluvisol and the Gleysol, respectively. We sampled the soils using an auger with one sample collected every 10 cm respecting the horizons boundaries.

In the Cambisol, we sampled the Ap- and B-horizons (Supplementary Table 1). The Ap-ploughing horizon (0 to 20 cm depth) has a silty texture and a dark brown colour (10YR3/1 according to the Munsell chart). The B-horizon (20 to 60 cm depth) has a silty texture and a yellowish-brown colour (10YR5/6). The ratio of 2-20 µm to 20-50 µm fractions is close to one indicating a parent material of eolian origin for the sampled horizons (Walter and Curmi, 1998), likely similar to the loess of northern and/or western Brittany (Haase et al., 2007; Le Calvez, 1979).

In the Albeluvisol, we sampled the Ap- and Bt-horizons (Supplementary Table 1). The Ap-horizon (0 to 20 cm depth) has a silty texture and a dark-brown to greyish colour (10YR3/2 to 10YR4/2). The Bt-horizon (20 to 60 cm depth) is silty with clay concentration increasing with depth. The ratio of 2-20 µm to 20-50 µm fractions in all the horizons is close
to 2 indicating that the soil profile is developed from the sedimentary Brioverian schist parent material (Walter and Curmi, 1998).

In the Gleysol, we sampled the Ap- and B/C-horizons (Supplementary Table 1). The Ap-horizon (0 to ~25 cm depth) is dark grey (10YR2.5/1) with a silty-sandy-clayey texture. The illuvial B/C-horizon (~30 to 80 cm depth) is brown to brown-dark yellow (7.5 to 10YR4/6) with silty to slightly clayey texture. The ratio of 2-20 µm to 20-50 µm fractions in all the horizons is close to 2 suggesting that the soil mainly developed from the sedimentary Brioverian schist parent material (Walter and Curmi, 1998).

The Albeluvisol and Gleysol contain redoximorphic features such as mottling and rust accumulations in root spaces. These features are of millimetric to centimetric size, of brown, yellowish to red or pale grey to greenish colour. They are sparse in the upper Ap horizon but frequently present in the deeper Bt and B/C horizons of the Albeluvisol and Gleysol.

2.2. Methods and analyses

Collected samples were dried at 40 °C and sieved to < 2 mm. Aliquots were taken for elemental and isotopic analyses.

2.2.1. Fe concentrations: bulk analyses and selective extractions

Total iron concentration of the bulk soils was determined on 0.25 g aliquots. Each aliquot was dissolved in a HF-HClO₄ mixture after calcination of the organic matter (450 °C). We also used two methods of selective extractions using reducing agents of increasing strength to obtain information about the major pools of iron in the soils studied: (1) the Tamm’s extraction in the dark and (2) the citrate-bicarbonate-dithionite (CBD)-extraction.

The Tamm’s reagent is a mixture of oxalic acid and ammonium oxalate (Tamm, 1922). The extraction was performed by shaking the sample-solution mixture over 4 hours, at 20 °C and in the absence of light with a solid/liquid ratio of 1.25 g/50 ml. This method allows
the extraction of weakly bound, poorly crystalline and organic-bound iron (Duchaufour and Souchier, 1966). For the extraction by CBD, the soil sample was exposed to the reactant mixture at 80°C with a solid/liquid ratio of 0.5g/25mL during 30 min. This method extracts the iron bound to oxides and hydroxides (hematite, goethite, lepidocrite) (Mehra and Jackson, 1960).

Bulk soil iron concentrations and iron concentration in the solutions from the partial extractions were analysed using an Inductively Coupled Plasma-Atomic Emission Spectrometer (LAS Arras).

Each extraction was done from an aliquot of the bulk soil. While the Tamm reagent extracts the weakly-bound Fe, the CBD extracts both the weakly bound and the oxide bound-Fe. We calculated the oxide-bound Fe concentration by subtracting the weakly-bound Fe measured by Tamm reagent extraction from the CBD extraction and the silicate-bound Fe concentration by subtracting the weakly-bound and the oxide-bound iron from the total iron concentration.

### 2.2.2. Fe isotope ratios measurements

Samples for the measurements of Fe isotope ratio were prepared and analysed at the Ecole normale supérieure de Lyon (ENS Lyon), France. An aliquot of each sample was ground using an agate mortar. Approximately 300 mg of sample powder was first treated with 30% H2O2 in order to eliminate the organic matter and then dissolved using a mixture of concentrated HF-HNO3-HCl acids, at ~130°C. Iron was separated and purified by anion exchange chromatography (AG MP1, 100-200 mesh, chloride form). Samples were loaded on the resin in 7N HCl-0.001% H2O2 and Fe was eluted with 2N HCl-0.001% H2O2 (Maréchal et al., 1999). As anion-resins could fractionate Fe isotopes, complete recovery is necessary. The yield was found to be better than 99%. For the iron isotopes ratios measurements, we used a high-resolution MC-ICP-MS (Nu Plasma 1700, Nu Instruments) at ENS Lyon, operated at a
mass-resolution (m/Δm) of 3000 ± 100, in dry plasma mode using the Nu DSN desolvation system. Samples were introduced by free aspiration in 0.05N sub-boiled distilled HNO₃ using a glass microconcentric nebulizer (uptake rate: 100 μl/min). We used a standard-bracketing approach with IRMM-014 as standard reference material to correct for instrumental mass bias using the exponential law (Albarède and Beard, 2004). Sample measurement solutions were diluted to match the concentration of the IRMM-014 standard within 15%, i.e. 150 to 300 ppb depending on the measurement session. The precision of the isotopic compositions (external reproducibility, 2σ) calculated on the basis of repeated measurement of the IRMM-014 standard were 0.11 and 0.17 ‰ (N= 324) for δ⁵⁶Fe and δ⁵⁷Fe, respectively. In a δ⁵⁷Fe vs. δ⁵⁶Fe diagram, all soil samples measurements plot along a line with a slope of 1.46 (Supplementary Fig. 1). This value is equal, within error margins, to the theoretical value of ln(M⁵⁷/M⁵⁴)/ln(M⁵⁶/M⁵⁴) = 1.487, indicating mass-dependent fractionation and no influence of isobaric interferences.

3. Results

The results for the three soil profiles are presented in figures 1 and 2 and in supplementary table 1. Classical soil parameters, such as granulometry, pH, and C and N concentrations, as well as Fe concentrations and isotopic compositions are indicated. The last two sets of results are presented in details in the following sections.

3.1. Fe concentration

Total iron concentration was found to be uniform along the Ap-B horizons of the Cambisol with only a slight increase at the bottom of the B horizon (Fig. 1). In the Albeluvisol and Gleysol, a sharp decrease (- 330 to 350%) of the total iron concentration in the surface horizons was observed with respect to the deep horizons (Fig. 1b). In the Bt
horizon of the Albeluvisol, the iron concentration is maximum around 25 cm depth and decreases slightly below. In the B/C horizon of the Gleysol, the iron concentration increases slightly with depth. In the surface horizons, the Cambisol located at the hilltop had the highest Fe concentration compared to those of the Albeluvisol and Gleysol. The Fe concentrations measured for the three soils were close to those obtained by X-ray fluorescence by Trolard et al. (2002) for nearby locations of the same soil.

Selective extractions showed that the contributions of the different Fe pools varied between the horizons and the soil types (Fig. 1c-e and supplementary Table 1). The oxide-bound Fe, corresponded to 39 to 81% of the total iron (Supplementary Fig. 2) and its vertical evolution in each soil mimicked that of the total Fe.

The weakly-bound, poorly crystalline iron pool corresponded to 1 to 18% of the total Fe (Supplementary Fig. 2). It was higher in the Cambisol than in the two other profiles. In the three soils, it increased progressively up to the surface horizon.

The silicate-bound Fe represented 18 to 45% of the total iron (Supplementary Fig. 2) and decreased gradually from depth to surface in the Cambisol. In the Albeluvisol, it increased progressively from a depth of 60 cm to reach a maximum around a depth of 20-30 cm, then showed a sharp decrease at the transition between the Bt and the Ap horizons around 20 cm. For the Gleysol, its maximum value was recorded in the deepest sample and decreased progressively to the top of the B/C horizon. A decrease was observed at the B/C – Ap transition. The Ap horizons have similar silicate-bound Fe concentrations in the three profiles.

### 3.2. Fe isotopic compositions

The iron isotopic composition in the studied soils varied from $\delta^{56}\text{Fe} = -0.15$ to 0.26‰ (Fig. 2 and supplementary Table 1). The three studied soils show differences in the iron isotopic compositions and variations with depth. The Cambisol profile showed variations of
δ^{56}Fe from 0.00 to only 0.15 ‰ indicating virtually no vertical fractionation along this profile. In contrast, the amplitude of the Fe isotopic variations was much larger than the external reproducibility within the two other profiles. In the Albeluvisol, δ^{56}Fe ranged from 0.00 to 0.26 ‰ with a decrease of the δ^{56}Fe values from the surface down to ca. 40 cm. In the Gleysol, δ^{56}Fe varies from -0.15 to 0.22 ‰ with values decreasing with depth. Notably a marked decrease (-0.19 ‰) was found between the surface Ap horizon and the deeper B/C horizon, corresponding to the observed increase in total Fe concentration (Fig. 1b) and variations in the relative proportions of the various Fe pools (Supplementary Fig. 2). In both the Albeluvisol and the Gleysol, the surface horizons were enriched in heavy Fe isotopes while the deep horizons were enriched in light isotopes (Fig. 2). The non-hydromorphic Cambisol exhibited the lowest δ^{56}Fe value and the highest Fe concentration in the surface horizon whereas it was the opposite for the hydromorphic Albeluvisol and Gleysol.

4. Discussion

Heterogeneities observed in soils can result from different pedogenetic processes and factors, such as the parent material and topography (Jenny, 1941) that contribute to the soil formation. The soils studied in this work can be differentiated by two main factors: the parent material and the water regime. The Cambisol developed on material of eolian origin under oxic conditions in a well-drained environment. In contrast, the other two soils formed on sedimentary Brioverian schists and remain waterlogged during a great part of the year and are thus subject to reduction processes. The evolution of the Fe behaviour with depth will thus be discussed separately, on the one hand in the Cambisol and on the other hand in the two waterlogged soils in an attempt to identify the mechanisms that control the Fe transfers.

4.1. Evolution of the Fe in the Cambisol
In the Cambisol, the amount of oxide-bound Fe was found to remain constant, whereas the amount of weakly-bound Fe increased and that of Si-bound Fe decreased from the base to the top of the soil profile (Fig. 1c-e). A question arises about whether an exchange could occur between these last two Fe pools and whether Fe remobilization could take place in this soil profile. The Cambisol profile is characterised by an overall homogeneous bulk Fe isotopic signature (Fig. 2): $\delta^{56}\text{Fe} = 0.00$ to 0.15‰. Two scenarios could explain this rather uniform isotopic composition: (1) Fe redistribution processes took place but did not fractionate the Fe isotopes or (2) soil evolution processes happened at the centimetric to decimetric scale, i.e. no major vertical Fe translocation occurred in this soil and Fe export out of this soil was extremely limited. In the deepest sample having the highest Si-bound Fe value, the total Fe concentration was 31.1 g kg$^{-1}$ and the $\delta^{56}\text{Fe}$ was 0.0‰. This concentration value is within the range of those of Brittany loess (22.0 ± 17.2 g kg$^{-1}$ (2σ), Gallet et al. (1998)). In addition, both the concentration and isotopic composition values are, within experimental error, close to those of worldwide loess: $[\text{Fe}_{\text{total}}] = 22.7 \pm 8.8$ g kg$^{-1}$ (Taylor et al., 1983) and $\delta^{56}\text{Fe} = 0.05 \pm 0.04$ ‰ (Beard et al., 2003b). Both observations are in agreement with the fact that the loess is the parent material of the Cambisol (see section 2.1). Mineral weathering by hydrolysis in the surface soil could have liberated Si-bound Fe as indicated by the lower values relative to the rest of the profile (Fig. 1e). The comparison between the average value of the Ap horizon and the deepest sample of the B horizon shows that there is a 4.7 g kg$^{-1}$ decrease of the total Fe concentration in the surface horizon (Fig. 1b). In addition, in the homogenous Ap-horizon, 4.4 g$_{\text{Fe}}$ kg$^{-1}$ is lost from the silicate compartment (Fig. 1e) while 2.2 g$_{\text{Fe}}$ kg$^{-1}$ is gained in the weakly-bound Fe pool (Fig. 1d). These observations show that part of the iron is removed from the profile. The loss in the Ap horizon corresponds, however, to a very small decrease of the total Fe concentration (ca. 15%) between the deepest sample and the two surface samples of the ploughing horizon.
There is also an in-situ transfer from the Si-bound to the weakly-bound iron pool representing about half of the mobilised Fe. To summarize, the redistribution of Fe in the Cambisol was found to be very limited and the net loss of Fe too small to generate significant Fe isotopes fractionation over the Cambisol profile.

4.2. Fe transport in the hydromorphic soils (Albeluvisol and Gleysol)

The oxide-bound Fe is the dominant pool of Fe in the hydromorphic soils investigated (Fig. 1c and supplementary Fig. 2a). There is a sharp decrease in the oxide-bound Fe concentration in the surface horizons in both soils concomitant with a decrease in total Fe concentration (Fig. 1b, c) suggesting that there is a net export of Fe from these surface horizons. This export can be explained by the presence of a water table that develops in the surface horizons during winter-spring and creates reducing conditions favourable to Fe remobilization. The similarity of both the total Fe concentrations (Fig. 1b) and the isotopic compositions (Fig. 2) in the surface horizons of these soils suggests that the Fe transport in these soils is due to the same process. Dissolution of Fe in the surface horizons would create (1) a decrease in the Fe concentration (Fig. 1b) and (2) a residual Fe pool enriched in heavy isotopes (Fig. 2) because light Fe isotopes are preferentially remobilized by dissolution. The isotopically light product of the dissolution reaction could be transported laterally out of the soil profiles along the topography or vertically to greater soil depth.

The iron isotopic composition of the bulk soil is mostly controlled by this process as shown by the strong negative correlation ($R^2 = 0.70$, $p < 0.01$, data normally distributed) between the $\delta^{56}$Fe of the bulk soil samples and the oxide-bound Fe concentration (Fig. 3). Accordingly, the dissolution of Fe-bearing minerals and transport of the free Fe (II) would be the processes that dominate the Fe movements in these hydromorphic soils.
By contrast, as aforementioned, there is no relationship between the oxide-bound Fe concentration and the $\delta^{56}$Fe value of the bulk soil samples in the Cambisol (Fig. 3). This confirms that the mechanism governing the evolution of Fe in the Cambisol is different from that involved in the Albeluvisol and Gleysol.

The Albeluvisol and Gleysol profiles below the Ap horizon are more difficult to interpret. Both soils developed from a sedimentary Brioverian schist parent material (see section 2.1) which is a highly heterogeneous material as shown, in particular, by the complex evolution of the Si-bound iron profiles. Thus, no mass balance calculations could be performed.

In the B/C horizon of the Gleysol, the observed overall variation of the Fe isotopic composition with depth is almost an inverse image of the variation of the Si-bound Fe with depth (Figs. 1e and 2). Indeed, there is a strong negative correlation between the $\delta^{56}$Fe and the Si-bound concentrations (Supplementary Fig. 3a). This feature could be interpreted as reflecting the imprint of the parental material on the isotopic composition of the Gleysol. Thus, the isotopic composition of the parental material appears to be the main factor that governs the Fe isotopic signature of the B/C horizon of the Gleysol. On the contrary, there is no such relationship in the Bt horizon of the Albeluvisol (Supplementary Fig. 3b). Thus, the parental material has little influence, if any, on the Fe isotopic composition of the Albeluvisol Bt profile.

In the Bt horizon of Albeluvisol, the two deepest samples are both enriched in Fe heavy isotopes and depleted in oxide-bound Fe relative to the corresponding Gleysol samples (Figs. 1c and 2). In addition, the Fe concentration profiles (Figs. 1b-e) could reflect a greater dissolution of Fe with increasing depth. Indeed, an underground water table is seasonally present in the Bt horizon of the Albeluvisol (see section 2.1; Davranche et al., 2011; Olivié-Lauquet et al., 2001; Pauwels et al., 1996). It results in reducing conditions which allow Fe
dissolution and the remobilisation of oxide-bound Fe from the flooded horizons of the Albeluvisol. This process favours the removal of light Fe isotopes and leaves a residual Fe pool enriched in heavy isotopes. This is in agreement with the observed negative correlation between the $\delta^{56}$Fe of the bulk soil samples and the oxide-bound Fe concentration (Fig. 3). The mobile, isotopically light Fe(II), could be transported down the topography. Thus, the main process that appears to drive the Fe isotopic composition in the Bt horizon of the Albeluvisol is dissolution due to the variation of the level of the underground water table.

Our results suggest that both, variation of the underground water level and topography-driven lateral water flows are involved in the translocation of Fe. The isotopically-light Fe removed from the Albeluvisol could be redeposited deeper within the Albeluvisol or in the Gleysol located down the slope or be evacuated from the toposequence. Due to the lack of data on the Fe concentration and isotopic composition on the parent material we have however no clear evidence for such redeposition. This study demonstrates that Fe isotopic compositions can evidence Fe transfers where mass balance calculations are not feasible and pinpoint the factors that control the distribution of Fe within a soil profile.

**4.3. Comparison to Fe isotopic compositions in worldwide soils**

We have compiled published Fe isotope composition of bulk horizons from different soil types (Table 1 and Fig. 4). The overall range for the bulk soil is $\delta^{56}$Fe = -0.62 to 0.72 ‰.

We have divided the soils into two groups on the basis of the range of their isotopic variations and we have tried to pinpoint the common characteristics for the soils in each group in an attempt to distinguish the factors that control the Fe isotopic fractionations. Group 1 is formed by the soils with small vertical Fe isotopic fractionation, i.e. $\Delta^{56}$Fe$_{profile} \leq 0.15$ ‰, with $\Delta^{56}$Fe$_{profile}$ being the difference between the highest and the lowest $\delta^{56}$Fe values in a given profile (Table 1 and Fig. 4). It regroups the Cambisols ($\Delta^{56}$Fe$_{profile} =$
0.08 to 0.15 ‰), the Ferralsols ($\Delta^{56}\text{Fe}_{\text{profile}} = 0.08$ to $0.14$ ‰) and the Haplic Gleysol ($\Delta^{56}\text{Fe}_{\text{profile}} = 0.11$ ‰).

Group 2 consists of soils with significant Fe isotopic fractionation along a vertical profile, such as the Podzols ($\Delta^{56}\text{Fe}_{\text{profile}} = 0.24$ to $0.77$ ‰), the Albeluvisol ($\Delta^{56}\text{Fe}_{\text{profile}} = 0.26$ ‰), the Gleysol ($\Delta^{56}\text{Fe}_{\text{profile}} = 0.41$ ‰), the Ultisol ($\Delta^{56}\text{Fe}_{\text{profile}} = 0.74$ ‰) and the Andosol ($\Delta^{56}\text{Fe}_{\text{profile}} = 0.30$ ‰).

Within group 1, the Cambisols and the Ferralsols develop under oxic and well-drained conditions with a dominant vertical water transfer. Yet, the absence of the Fe isotopic fractionation along the profile suggests that either the water circulation has no effect on the oxidation state of Fe as suggested for Ferralsols by Poitrasson et al. (2008), or that mineral dissolution is not associated with long-distance transfer of Fe and that the released Fe precipitates to form secondary minerals without transport as suggested by Wiederhold et al. (2007a). It is worth noticing that the temporal evolution of the soils does not appear to influence the isotopic signature; indeed, the Cambisols were young, poorly developed soils, whereas the Ferralsols result from a long-term evolution, up to several millions of years (Poitrasson et al., 2008).

The Haplic Gleysol and the Stagnic Cambisol show a range of $\delta^{56}\text{Fe}$ variations similar to those of the other profiles of group 1 ($\Delta^{56}\text{Fe}_{\text{profile}} < 0.15$ ‰), however, they developed under seasonally or permanently water-saturated, anoxic conditions (Wiederhold et al., 2007b). While reductive Fe mobilization under anoxic conditions was inferred to occur in both soils (Wiederhold et al., 2007b), the absence of large isotopic variations was suggested to indicate that the transport of Fe within each profile was spatially limited and, hence, that most of the Fe would be transformed at the centimetric to decimetric scale.

Group 2 of soil profiles correspond to $\Delta^{56}\text{Fe}_{\text{profile}} > 0.15$ ‰ (Table 1 and Fig 4). The Podzols developed under well-aerated conditions with vertical water transfer (Emmanuel et
al., 2005; Wiederhold et al., 2007a). Unlike Cambisols from group 1 that evolved in similar settings, the Podzols were characterised by acidic pH. Under these pH conditions, Fe(III) could be dissolved and the released isotopically light Fe could vertically be translocated by organic matter. At depth, isotopically light Fe could precipitate. These processes would result in significant Fe isotopes fractionation.

The other soils of group 2 – the Albeluvisol, the Gleysol and the Ultisol – developed under poorly-drained to water-saturated conditions and were subject to variations of the redox conditions due to changes in water-saturation states. In all cases, the reductive dissolution of Fe would affect preferentially the light isotopes which become mobile. Hence, the residual material would become isotopically heavier, while the levels where Fe accumulation occurred would become enriched in isotopically light Fe.

The Andosol has an intermediate behaviour between the Podzols and the water-saturated soils. It formed along volcanic slope in Hawaii with a mean annual precipitation of 3500 mm/year. Water-saturated (Thompson et al., 2007) and low oxygen availability (Schuur et al., 2001) conditions would take place during the periods of higher precipitation. Iron isotope fractionation would result from the reductive dissolution of Fe, and $\delta^{56}\text{Fe}$ values would increase with increasing removal of Fe.

To conclude, iron isotopes represent a great tool for studying the mechanisms and scales (centimetric- to toposequence-) of Fe transfer in soils and distinguishing between various mechanisms of soil evolution. For these reasons, they should be used in combination with pedological and geochemical characterisation as well as an appropriate characterisation of the geographical and environmental contexts.

5. Conclusions
Iron isotopes profiles in soils have been shown to be a tool of great potential to better understand the fate of Fe in soils and the evolution of soils. In this study, iron isotopic variations have helped us to distinguish between various mechanisms of Fe transport within a soil profile and along a toposequence.

For the Cambisol, Fe isotopic fractionations were small and Fe transfers were limited. The mechanisms corresponding to these observations were: (1) centimetric to decametric-scale exchanges between iron pools and (2) lateral transport of Fe down the slope; the latter representing only a small proportion of the total Fe.

The bulk Fe isotopic composition in the hydromorphic soils (Albeluvisol and Gleysol) was found to be dominated by the behaviour of the oxide-bound Fe. In the surface horizon, the main mechanism of Fe transport in these soils is a strong Fe dissolution and transport of isotopically light Fe out of these horizons. Even if Fe could be transferred within each profile, it would mainly be transported laterally along the slope. The fluctuations of the underground water level could also induced the translocation of Fe after dissolution of the oxides during the periods of reducing conditions corresponding to the high level stands. Iron isotopic compositions have allowed us to show that this phenomenon is dominant in the Bt horizon of the Albeluvisol. Finally, Fe isotopic composition of the B/C horizon of the Gleysol is dominated by the signature of the parental material.

Iron isotopic compositions have allowed us to (1) distinguish between the in-situ processes (centimetric- to decimetric-scale) and the processes inducing Fe transfer at the profile- and toposequence-scale, and (2) identify the mechanisms or factors that control the Fe distribution in soils. Consequently, determining the Fe isotopic compositions represents a powerful tool for studying the fate of Fe in soil systems when no sufficient data are available for full mass balance calculations.
More generally, the comparison we made of Fe isotopes profiles in various soils of the world shows that Fe isotope ratio variations along a soil profile are linked to the evolution processes that have affected the soil. Hence, Fe isotopic compositions in soils can help us discriminate between various mechanisms of soil evolution when they are used in combination with pedological, geochemical, geographical, and environmental characterisations.
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FigureCaptions

**Fig. 1.** Variations of the soil characteristics with depth in the three studied soil profiles. (a) clay concentration, (b) total Fe concentration, (c) oxide-bound Fe extracted with CBD, (d) weakly-bound Fe extracted with Tamm’s reagent, and (e) calculated Si-bound Fe (see the Material and Methods section for details). The vertical dashed line in (b) represents the average Fe concentration (50.5 g kg\(^{-1}\)) for the Bt and B/C horizons of the Albeluvisol and Gleysol, (see section 4.2 for details).

**Fig. 2.** Variations of the iron isotopic compositions (\(\delta^{56}\text{Fe}\)) with depth in the three studied soil profiles.

**Fig. 3.** Relationship (black line) between the bulk iron isotopic compositions and the oxide-bound Fe concentration in the Albeluvisol and the Gleysol. There is no relationship for the Cambisol.

**Fig. 4.** Global patterns of bulk Fe isotope ratio variations in the Naizin soils and other soils worldwide. Two behaviours are distinguished: soil profiles with limited \(\delta^{56}\text{Fe}\) variations (group 1), and soils in which significant \(\delta^{56}\text{Fe}\) variations were measured (group 2). The details for each soil are indicated in Table 1.
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Figure 1
Figure 2
$y = -0.01x + 0.25$

$R^2 = 0.69$

Figure 3
Figure 4

GROUP 1: $\Delta^{56}\text{Fe}_{\text{profile}} < 0.15\%e$

Well-drained conditions, vertical water transfer

GROUP 2: $\Delta^{56}\text{Fe}_{\text{profile}} > 0.15\%e$

Poorly-drained conditions, lateral water transfer or no transfer
<table>
<thead>
<tr>
<th>Soil type</th>
<th>Location</th>
<th>Bedrock</th>
<th>Vegetation cover</th>
<th>pH</th>
<th>$^{56}\text{Fe}$ range,‰</th>
<th>$^{56}\text{Fe}_{\text{profile}}$,‰</th>
<th>N</th>
<th>[Fe], g/kg</th>
<th>Reference</th>
<th>Differen-</th>
<th>Redox</th>
<th>Vertical transfer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cambisol</td>
<td>Kervidy-Naizin, Brittany, NW France (48°00'N, 2°30'E)</td>
<td>eolian deposits</td>
<td>crop</td>
<td>6.0 to 6.5</td>
<td>0.00 to 0.15</td>
<td>0.15</td>
<td>6</td>
<td>25.8 to 31.1</td>
<td>this study</td>
<td>-</td>
<td>-</td>
<td>+</td>
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<tr>
<td>Cambisol</td>
<td>Immendingen, Hegau region, SW Germany (47°54'N, 8°44'E)</td>
<td>basaltic tuff (Hewenegg tertiary volcanic complex)</td>
<td>beech trees</td>
<td>5.5 to 5.9</td>
<td>0.11 to 0.21</td>
<td>0.10</td>
<td>4</td>
<td>111.4 to 115.0</td>
<td>Wiederhold et al. (2007a)</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Cambisol</td>
<td>Shaar Hagay area, Judean Mountains, Israel</td>
<td>limestone and dolomite</td>
<td>semi-arid vegetation</td>
<td>6.8 to 7.9</td>
<td>-0.21 to -0.13</td>
<td>0.08</td>
<td>3</td>
<td>43.8 to 54.1</td>
<td>Emmanuel et al. (2009)</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Cambisol</td>
<td>Ratz, Canton Zurich, Switzerland (47°37'N, 8°44'E)</td>
<td>glacial moraine deposit + eolian silt</td>
<td>spruce trees</td>
<td>3.4 to 4.1</td>
<td>0.02 to 0.16</td>
<td>0.14</td>
<td>6</td>
<td>18.0 to 33.0</td>
<td>Wiederhold et al. (2007b)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ferralsol</td>
<td>Laterite, 120 km SE of Yaoundé, Cameroon (3°10'N - 11°58'E)</td>
<td>granodiorite</td>
<td>semi-deciduous rainforest</td>
<td>no data</td>
<td>0.08 to 0.15</td>
<td>0.08</td>
<td>7</td>
<td>11.3 to 193.7</td>
<td>Poitrasson et al. (2008)</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Ferralsol</td>
<td>Laterite, 120 km SE of Yaoundé, Cameroon (3°10'N - 11°58'E)</td>
<td>granodiorite</td>
<td>semi-deciduous rainforest</td>
<td>no data</td>
<td>0.02 to 0.16</td>
<td>0.14</td>
<td>6</td>
<td>30.5 to 148.3</td>
<td>Poitrasson et al. (2008)</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Podzol (P1)</td>
<td>Flaesheim/ Westphalia, NW Germany (51°430 N, 7°120 E)</td>
<td>eolian sand deposit (Pleistocene)</td>
<td>pine trees</td>
<td>3.1 to 4.0</td>
<td>-0.29 to 0.47</td>
<td>0.77</td>
<td>10</td>
<td>1.9 to 19.4</td>
<td>Wiederhold et al. (2007a)</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Podzol (P2)</td>
<td>Klosterreichenbach/Black Forest, SW Germany (48°310 N, 8°240 E)</td>
<td>hematitic sandstone (Buntsandstein, Trias) reworked by periglacial processes</td>
<td>spruce and pine trees</td>
<td>2.7 to 4.6</td>
<td>-0.02 to 0.51</td>
<td>0.53</td>
<td>10</td>
<td>1.9 to 14.2</td>
<td>Wiederhold et al. (2007a)</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Podzol (P3)</td>
<td>Naetàlm site, Krudne hory Mountains, NW Czech Republic</td>
<td>quartzite and biotite - sillimanite gneiss</td>
<td>forest</td>
<td>3.1 to 4.1</td>
<td>-0.31 to -0.07</td>
<td>0.24</td>
<td>6</td>
<td>13.2 to 41.5</td>
<td>Emmanuel et al. (2005)</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Gleysol (G1)</td>
<td>Tettang, Upper Swabia, S Germany (47°39'N, 9°33'E)</td>
<td>carbonates - sandy sediments (Pleistocene) + a layer of loamy sediments</td>
<td>deciduous trees</td>
<td>5.9 to 7.7</td>
<td>0.01 to 0.12</td>
<td>0.11</td>
<td>6</td>
<td>13.0 to 26.7</td>
<td>Wiederhold et al. (2007b)</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Gleysol (G2)</td>
<td>Kervidy-Naizin, Brittany, NW France (48°00'N, 2°30'E)</td>
<td>sedimentary Brioverian schist (older than 530 Ma)</td>
<td>poplar trees</td>
<td>5.4 to 6.3</td>
<td>-0.15 to 0.26</td>
<td>0.41</td>
<td>7</td>
<td>10.6 to 63.4</td>
<td>this study</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Albeluvisol (Alb)</td>
<td>Kervidy-Naizin, Brittany, NW France (48°00'N, 2°30'E)</td>
<td>sedimentary Brioverian schist (older than 530 Ma)</td>
<td>poplar trees</td>
<td>5.4 to 5.6</td>
<td>0.00 to 0.26</td>
<td>0.26</td>
<td>6</td>
<td>11.7 to 52.7</td>
<td>this study</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Andosol (?) (And)</td>
<td>3500 mm rain site, Maui, Hawaii, USA (20°48'N, 156'10'W)</td>
<td>basalts</td>
<td>montane rainforest</td>
<td>no data</td>
<td>0.15 to 0.45</td>
<td>0.30</td>
<td>4</td>
<td>54.3 to 97</td>
<td>Thompson et al. (2007)</td>
<td>+</td>
<td>+</td>
<td>?</td>
</tr>
<tr>
<td>Ultisol (?) (UR)</td>
<td>Ferncreek series, Mendocino, California, USA</td>
<td>lithified marine sediments (Franciscan complex, late Cretaceous - early Tertiary)</td>
<td>Bishop pine ± Douglas fir ± redwood</td>
<td>3.9 to 4.9</td>
<td>-0.61 to 0.13</td>
<td>0.74</td>
<td>3</td>
<td>no data</td>
<td>Fantle and DePaolo (2004)</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
</tbody>
</table>

**Table 1**: Compilation of bulk Fe isotopic compositions in worldwide soil profiles and characteristics of these soils.