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Geochemical modeling of Fe(II) binding to humic and fulvic acids

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ABSTRACT

The complexation of Fe(II) with organic matter (OM) and especially with humic acids (HA) remains poorly characterized in the literature. In this study, batch experiments were conducted on a pH range varying from 1.95 to 9.90 to study HA-mediated Fe(II) binding. The results showed that high amounts of Fe(II) are complexed with HA depending on the pH. Experimental data were used to determine a new set of binding parameters by coupling PHREEPLOT and PHREEQC-Model VI. The new binding parameters ($\log K_{MA} = 2.19 \pm 0.16$, $\log K_{MB} = 4.46 \pm 0.47$ and $\Delta LK_2 = 3.90 \pm 1.30$) were validated using the LFER (linear free energy relationship) method and published adsorption data between Fe(II) and Suwannee River fulvic acid (SRFA) (Rose and Waite, 2003). They were then put in PHREEQC-Model VI to determine the distribution of Fe(II) onto HA functional groups. It was shown that Fe(II) forms mainly bidentate complexes, some tridentate complexes and only a few monodentate complexes with HA. Moreover, Fe(II) is mainly adsorbed onto carboxylic groups at acidic and neutral pH, whereas carboxy-phenolic and phenolic groups play a major role at basic pH. The major species adsorbed onto HA functional groups is Fe^{2+} ; $Fe(OH)^+$ appears at basic pH (from pH 8.13 to 9.9). The occurrence of OM and the resulting HA-mediated binding of Fe(II) can therefore influence Fe(II) speciation and bioavailability in peatlands and wetlands, where

seasonal anaerobic conditions prevail. Furthermore, the formation of a cationic bridge and/or the dissolution of Fe(III)-(oxy)hydroxides by the formation of Fe(II)-OM complexes can influence the speciation of other trace metals and contaminants such as As.

1. Introduction

Natural organic matter (NOM) derives from the biological and chemical degradation/transformation of plant and animal residues (Piccolo, 1996). The most refractory and reactive fraction with regards to metal binding is composed of humic substances (HS). Humic substances can be divided into three operational fractions: humic acids (HA) soluble under alkaline conditions, fulvic acids (FA) soluble over the entire pH range and humins, the insoluble fraction. Humic substances, and especially HA and FA, are considered to exert a major control on metal mobility and bioavailability in the environment (Sposito, 1986; Buffle et al., 1998; Tipping, 2002). They are renowned for their ability to bind a large range of metals and metalloids including possibly toxic elements such as REE, Al(III), Pb(II), Ca(II), Mn(II), Mg(II), Fe(III) and smaller amounts of As(V) (Thanabalasingam et al., 1986; Redman et al., 2002; Lin et al., 2004; Buschmann et al., 2004; Bauer et al., 2006; Ritter et al., 2006; Kar et al., 2011), As(III) (Thanabalasingam et al., 1986; Warwick et al., 2005; Buschmann et al., 2006; Liu et al., 2010), Sb(V) (Pilarski et al., 1995; Tighe et al., 2005; Filella et al., 2012) and Sb(III) (Pilarski et al., 1995; Buschmann et al., 2004; Tella et al., 2009; Filella et al., 2012). The high binding capacity of HS is usually attributed to the large surface density of oxygen-containing functional groups (carboxylic, phenolic, carbonyl) and, to a lesser extent, nitrogen- or sulfur-containing functional groups (Evangelou et al., 2001). The binding ability of metals to HS has been intensively studied, notably through experimental and modeling studies (Fukushima et al., 1992; 1996; Town et al., 1993; Mota et al., 1994; Pinheiro et al., 1994; Tipping, 1998; Christl et al., 2000; Milne et al., 2003; Nuzzo et al., 2013). However, whereas HS binding with Fe(III) and Fe(III)-oxyhydroxides has been intensively studied (Weber et al., 2006; Van Schaik et al., 2008; Morris et al., 2012) only very few works have been dedicated

to the understanding of Fe(II)-HS complexation mechanisms (Schnitzer et al., 1966; Van Dijk, 1971; Rose and Waite, 2003; Yamamoto et al., 2010).

A better understanding and the quantification of Fe(II)-HS binding is crucial in the case of reduced waters, such as those occurring in peatlands, wetlands or anoxic sediments. As iron(II), the reduced species of Fe is much more soluble than Fe(III), Fe(II) concentrations are generally high (i.e. several mg L^{-1}) in these waters (Ponnamperuma, 1972; Buffle et al., 1989; Davison, 1993; Dia et al., 2000; Olivié-Lauquet et al., 2001). In such environments, where soils or sediments are periodically flooded and water-saturated, the organic matter (OM) is slightly degraded. High content of soluble, colloidal or particulate OM, and notably HS, are therefore encountered in these soils and soil solutions. The combination of these two features along with the high metal binding capacity of HS suggests that HS could be an important parameter in these waters, controlling not only the solubility, mobility and bioavailability of Fe(II), but also the kinetics of Fe(II) oxidation and the type of Fe(III) oxides formed during reoxidation (Pédrot et al., 2011).

Another more indirect reason to study the binding of Fe(II) by HS is that this binding could have major implications regarding groundwater contamination by As. Recent studies have shown that reduced, organic-rich sediments from flood plains could be an important source of As for underlying aquifers (Harvey et al., 2006; Kocar et al., 2008; Polizzotto et al., 2008; Fendorf et al., 2010). It has been shown in an oxic environment that Fe(III), through the formation of ternary complexes with HS, allows the binding of As(V) as an oxyanion to OM (Sharma et al., 2010; Mikutta et al., 2011). In this type of organic-rich environment, As(V) sorption onto OM seems to be a controlling factor for As speciation in contrast with previous studies which suggested that Fe-oxyhydroxides control the fate of As. Is it possible that similar complexes are formed between dissolved organic matter (DOM) and As(III) and As(V) in reduced, organic-rich waters, using Fe(II) as a cationic bridge between DOM and As species? Do these complexes enhance the mobility of As in flood plain sediments and to what extent are they involved in the contamination of underlying aquifers by As?

To better understand the potential role of HS with regards to Fe(II) and As mobility in reduced waters, a first step consists of describing and quantifying Fe(II)-HS binding. However, very few studies have been conducted to date with this purpose. It is necessary to identify which binding mechanisms are the prevailing ones and to estimate the stability constants of Fe(II) with HA and/or FA. Several attempts have been made to do this, but their results are contradictory. Schnitzer and Skinner (1966) established that Fe(II)-FA binding increases with increasing FA concentrations (from 0 to 1 g L⁻¹) and pH (from 3.5 to 5) with a maximum complexation at 57% of Fe(II) for pH = 3.5, [Fe(II)]_{tot} = 100 mg L⁻¹ and DOC = 553 mg L⁻¹. Van Dijk (1971) proposed from titration experiments that HA complexes Fe(II) through the formation of bidentate complexes. More recently, Rose and Waite (2003), who studied the kinetics of Fe(II) binding with 12 different NOMs in coastal waters, showed that Fe(II) binding with soil OM differs from one OM to another. Jackson et al. (2012) and Miller et al. (2009; 2012) studied the ability of OM to delay Fe(II) oxidation in an aerobic environment. Miller et al. (2012) proposed two mechanisms for Fe(II) oxidation in the presence of OM: first, a one-ligand model with two different oxidation mechanisms; second, they proposed a two-ligand model, where Fe(II) is first complexed with an OM strong binding site, and then with an OM weak binding site. Few datasets are therefore available concerning Fe(II)-HS binding.

These limited datasets could not be used to perform extensive modeling studies on Fe(II)-HS binding. Based on only one dataset (Van Dijk, 1971), Tipping (1998) reported the following Fe(II)-HA binding parameters: log K_{MA} = 1.28 and ΔLK₂ = 0.81 using Model VI. For the NICA-Donnan model, Milne et al. (2003) did not use Van Dijk's (1971) dataset because this dataset corresponds to pH data. They simply estimated the Fe(II)-HA binding parameters using the LFERs (linear free energy relationship) between their binding parameters and log K_{OH}, the first hydrolysis constant. Concerning Fe(II)-FA complexation, they used Schnitzer and Skinner's (1966) experimental datasets to determine their values for the binding parameters.

The aims of this study were to describe and quantify Fe(II)-HA complexation. Batch experiments involving complexation between Fe(II) and Leonardite HA were performed. This

new experimental dataset was obtained through the coupling of an experimental approach combining ultrafiltration and Inductively Coupled Plasma Mass Spectrometry (ICP-MS) measurements. PHREEPLOT-PHREEQC-Model VI was then used to model the binding of Fe(II) to HS using our new dataset and to determine the binding parameters of Fe(II). A validation of these values for the binding parameters was then performed using the LFER method and by reproducing published data using PHREEQC-Model VI. Speciation on the different sites was then determined, providing complexation mechanisms of Fe(II) to HA.

2. Experimental, analytical and modeling methods

2.1. Reagents and materials

All aqueous solutions were prepared with analytical grade Milli-Q water (Millipore). The Fe(II) stock solutions were prepared with iron chloride tetrahydrate ($\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$) from Acros Organics. NaOH, HCl and HNO_3 , all sub-boiling ultrapure grade, came from Fisher Chemical, Merck and VWR, respectively. Ammonium acetate, hydroxyammonium chloride and dimethyl-2,9 phenanthroline-1,10 chlorhydrate were obtained from Fisher Scientific, Merck and VWR Prolabo, respectively.

The HA used was the standard HA Leonardite from the International Humic Substance Society (IHSS). Prior to the experiments, HA molecules < 10 kDa were removed using a Labscale TFF system equipped with a Pellicon XL membrane (PGCGC10, MilliporeTM). After acidic digestion, blank Fe, Mn and Mg concentrations occurring in HA were determined by ICP-MS. The average concentrations were $206.6 \mu\text{g L}^{-1}$, $1 \mu\text{g L}^{-1}$ and $13.4 \mu\text{g L}^{-1}$ for Fe, Mn and Mg, respectively.

All materials were soaked in 10% HNO_3 and then rinsed with deionized water twice overnight. All experiments were conducted in a Jacomex isolator glove box (< 10 ppm of O_2) to prevent the oxidation of Fe(II) to Fe(III).

2.2. Setup of the binding experiments

A standard batch equilibrium technique was used. Three series of Fe(II)-HA complexation experiments were conducted in triplicate. Firstly, the pH was monitored and kept constant during 24 h with a multi-parameter Consort C830 analyzer combined with an electrode from Bioblock Scientific (combined Mettler InLab electrode). Calibrations were performed with WTW standard solutions (pH = 4.01 and 7.00 at 25°C). The accuracy of the pH measurements is ± 0.05 pH units. An isotherm adsorption experiment was carried out relative to the increasing Fe(II) concentration (0.61 to 8.55 mg L⁻¹). The average concentration of dissolved organic carbon (DOC) was 48.9 mg L⁻¹. The pH was fixed at 5.9 with ultrapure HCl and NaOH. Secondly, a pH sorption edge experiment was performed over a pH range from 1.95 to 9.90 with DOC and Fe(II) concentrations of 48.8 and 3.03 mg L⁻¹, respectively. Finally, a pH sorption-edge experiment was carried out over a pH range from 2.95 to 8.89 with DOC and Fe(II) concentrations of 76.5 mg L⁻¹ and 3.15 mg L⁻¹, respectively. The [Fe(II)]_{tot}, pH and DOC concentrations used in these experiments are representative of the concentrations that can be found in wetland waters (Ponnamperuma, 1972; Reddy and Patrick, 1977; Dia et al., 2000; Olivié-Lauquet et al., 2001). The ionic strength of all experiments was fixed at 0.05 M with NaCl electrolyte solution. Experimental solutions were stirred for 24 h to reach equilibrium. At equilibrium, 15 mL of solution was sampled and ultrafiltrated at 5 kDa (Vivaspin VS15RH12, Sartorius) by centrifugation at 2970 g for 30 min. under N₂ atmosphere. Ultracentrifugation cells were previously washed with 0.15 N HCl and Milli-Q water to obtain a DOC concentration below 1 mg L⁻¹ in the ultrafiltrate.

2.3. Chemical analyses

Dissolved organic carbon (DOC) concentrations were determined using an organic carbon analyzer (Shimadzu TOC-V CSH). The accuracy of the DOC measurements was estimated at $\pm 5\%$ for all samples using a standard solution of potassium hydrogen phthalate. Iron concentrations were determined by ICP-MS using an Agilent Technologies 7700x at

Rennes 1 University. The samples were previously digested twice with 14.6 N HNO₃ at 90°C, evaporated to complete dryness and then resolubilized with HNO₃ at 0.37 mol L⁻¹ to avoid any interferences with DOC during the analysis. ICP-MS analyses were carried out introducing He gas into collision cell to suppress any interference from Ar. The iron interference (⁴⁰Ar¹⁶O/⁵⁶Fe) was properly reduced by using He gas into collision cell to reach a low detection limit for Fe analysis (LD Fe: 0.07 µg L⁻¹) (Instrumental and data acquisition parameters can be found in the supplementary information). Quantitative analyses were performed using a conventional external calibration procedure (7 external standard multi-element solutions - Inorganic Venture, USA). A mixed solution of rhodium-rhenium at a concentration level of 300 ppb was injected on-line with the sample in the nebulizer. This solution was used as an internal standard for all measured samples to correct instrumental drift and matrix effects. Calibration curves were calculated from the intensity ratios between the internal standard and the analyzed elements. A SLRS-5 water standard was used to check the accuracy of the measurement procedure, and the instrumental error on the Fe analysis is < 5%. Chemical blanks of Fe were below the detection limit (0.07 µg L⁻¹), and were thus negligible.

Concentrations of Fe(II) in the ultrafiltrate ([Fe(II)]_{UF}) were determined with the 1,10-phenantroline colorimetric method (AFNOR, 1982) at 510 nm using a UV-visible spectrophotometer (UV/VIS Spectrometer "Lambda 25" from Perkin Elmer). Total Fe concentrations in the mixed HA-Fe(II) solutions were only measured by ICP-MS using the procedure described above. Indeed, the absorbance of Leonardite HA measured at 50 mg L⁻¹ of DOC is high (more than twice the absorbance of Fe(II)). The error of the [Fe(II)]_{UF} and [Fe(II)]_{tot} measurements was estimated at less than 5% above a concentration of Fe(II) of 0.85 mg L⁻¹ (7% for a concentration of Fe(II) of 0.6 mg L⁻¹). Iron (II) concentrations in the ultrafiltrates were assumed to be inorganic Fe(II), whereas Fe(II) bound to HA (Fe(II)-HA) was considered to be the retentate fraction > 5 kDa. The fraction of Fe(II) bound to HA was calculated as [Fe(II)-AH] = [Fe(II)]_{tot} - [Fe(II)]_{UF}, with [Fe(II)]_{tot} representing the Fe content of

the mixed HA-Fe(II) solutions prior to ultrafiltration and $[\text{Fe(II)}]_{\text{UF}}$ the Fe concentration determined in the ultrafiltrate by ICP-MS.

2.4. Determination of the PHREEQC-Model VI binding parameters

2.4.1. PHREEQC-Model VI

In this study, Model VI was coupled with PHREEQC as has been previously done (Marsac et al., 2012). It was used to calculate the partitioning of ions between the various complexing sites available on HS. PHREEQC-Model VI assumes that the complexation of ions by HS occurs through eight discrete sites: four weak sites, named A sites (usually assumed to be carboxylic groups), and four strong sites, named B sites (usually assumed to be phenolic groups). The abundance of type A and B sites are respectively named n_A (mol g^{-1}) and $n_B = 0.5 * n_A$ (mol g^{-1}). The intrinsic proton dissociation constants for type A and B sites and their distribution term are pK_A , pK_B , ΔpK_A and ΔpK_B , respectively. The fractions of proton sites that can make bidentate sites and tridentate sites are named f_B and f_T , respectively. Ion adsorption by humic and fulvic substances is described by a specific complexation parameter $\log K_{MA}$ and $\log K_{MB}$ for carboxylic and phenolic sites, respectively. The abundance of type A and B sites, their distribution term and the fraction of sites that can form bidentates and tridentates sites differ from HA to FA. The values of these constants are presented in Table 1 (from Tipping, 2002).

Tipping (1998) established a linear relationship between $\log K_{MA}$ and $\log K_{MB}$: $\log K_{MB} = 3.39 * \log K_{MA} - 1.15$ ($R^2 = 0.80$) to constrain the number of parameters set. This was deleted in PHREEQC-Model VI by Marsac et al. (2011), who determined the binding parameters of the rare earth elements (REE) with HA. The interaction between one site and one ion is characterized by the complexation constant $\log K$. These eight sites can therefore form monodentate, bidentate or tridentate complexes with a given ion. The $\log K$ values are calculated from (1) $\log K_{MA}$ and ΔLK_{1a} for a monodentate carboxylic site, (2) $\log K_{MB}$ and ΔLK_{1b} for a monodentate phenolic site, (3) $\log K$ of the two monodentate sites for a weak

bidentate site, (4) $\log K$ of the two monodentate sites and $1 \cdot \Delta LK_2$ (9% of the sites) and $2 \cdot \Delta LK_2$ (0.9% of the sites) for a strong bidentate site, (5) $\log K$ of the three monodentate sites for a weak tridentate site and (6) $\log K$ of the three monodentate sites and $1.5 \cdot \Delta LK_2$ (9% of the sites) and $3 \cdot \Delta LK_2$ (0.9% of the sites) for a strong tridentate site. Eighty binding equations are then defined. An electrical double layer is involved in the electrostatic interaction. The thickness of the electrical double layer corresponds to $1/\kappa$, where κ is the Debye-Hückel parameter. The ion distribution between the diffuse layer and the solution volume is calculated by a simple Donnan model.

2.4.2. PHREEPLOT

PHREEPLOT is a software used to create graphical output and to fit data using PHREEQC. The 80 complexation equilibria defined in PHREEQC-Model VI for HA and FA were added in the "minteq.v4" database. The humic and fulvic acids were defined as SOLUTION_MASTER_SPECIES, SOLUTION_SPECIES and PHASES. The 160 ($80 \cdot 2$) types of sites defined in PHREEQC-Model VI were added as SURFACE_MASTER_SPECIES in the "minteq.v4" database. The aim of the modeling process was to determine the value of $\log K_{MA}$ and $\log K_{MB}$ for Fe(II) by fitting the concentration of Fe(II) bound to HA from experimental data with those calculated by PHREEPLOT. A nonlinear relationship between $\log K_{MA}$ and $\log K_{MB}$ was used as previously performed by Marsac et al. (2011). The specific PHREEQC-Model VI binding parameters, namely $\log K_{MA}$, $\log K_{MB}$, ΔLK_{2C} and ΔLK_{2P} , were defined in PHREEPLOT as fitting parameters. Their values were determined by extrapolation of the present experimental Fe(II)-HA dataset. Binding parameters were optimized using the weighted sum of squares of the residuals. The stability constants of all sites defined in PHREEQC-Model VI were calculated from $\log K_{MA}$ and $\log K_{MB}$. The reactions necessary for PHREEPLOT calculation were added in the SURFACE_SPECIES section of PHREEPLOT using a different name for

each constant. The 80 equations defined in PHREEQC-Model VI were added using their SURFACE_SPECIES nomenclature in the PHREEPLOT input as "numericTags".

2.5. LFER linear free energy relationship

The PHREEQC-Model VI type A sites represent carboxylic groups of HA or FA. Acetic acid (CH_3COOH) can be used as a molecular model of HA carboxylic groups. Pourret et al. (2007) determined a linear relationship (LFER) between $\log K_{\text{MA}}$ and $\log K(\text{M-AA})$, the stability constant of a metal M with acetic acid:

$$\log K_{\text{MA}} = 1.03 * \log K(\text{M-AA}) - 0.43 ; \quad R^2 = 0.80 \quad (1)$$

Another LFER between $\log K_{\text{MA}}$ and the first hydrolysis constant of a metal M $\log K(\text{M-OH})$ has been suggested by the same authors (Pourret et al., 2007):

$$\log K_{\text{MA}} = 0.24 * \log K(\text{M-OH}) + 0.32 ; \quad R^2 = 0.78 \quad (2)$$

Marsac et al. (2011) showed that the linear relationship between $\log K_{\text{MA}}$ and $\log K_{\text{MB}}$ established by Tipping (1998) could not account for the binding of REE with HA. These authors estimated $\log K_{\text{MB}}$ using the LFER method based on $\log K_{\text{MB}}$ and the metal catechol stability constant ($\log K(\text{M-catechol})$):

$$\log K_{\text{MB}} = 0,37 * \log K(\text{M-catechol}) + 0,86 ; \quad (R^2 = 0.95) \quad (3)$$

3. Results

3.1. Experimental results

$\text{Fe(II)}_{\text{UF}}$ and $\text{Fe(II)}_{\text{tot}}$ concentrations (data presented in the supplementary information) determined using the phenanthroline method showed that no oxidation of Fe(II) occurred during any of the experiments. The quantification limit (QL) of the phenanthroline method is 0.5 mg L^{-1} of $\text{Fe(II)}_{\text{UF}}$ which is x5000 the QL of the ICP-MS method ($0.12 \text{ } \mu\text{g L}^{-1}$). The iron(II) concentrations used in this work were therefore those determined by ICP-MS.

3.1.1. Adsorption isotherm

Fig. 1a presents $\log [\text{Fe(II)}\text{-HA}]$ according to $\log [\text{Fe(II)}]_{\text{UF}}$. At low $[\text{Fe(II)}]_{\text{UF}}$ concentrations ($\log [\text{Fe(II)}]_{\text{UF}} = -6.2$), $\log [\text{Fe(II)}\text{-HA}]$ was approximately -5.0 and increased progressively to reach -4.28 for the maximum $\log [\text{Fe(II)}]_{\text{UF}}$ value used ($\log [\text{Fe(II)}]_{\text{UF}} = -4.00$). The lack of any plateau suggests that the saturation of HA complexation sites was not reached.

3.1.2. pH sorption edge

pH sorption edge experiments are very useful to better constrain the values of the binding parameters. Indeed, at acidic pH, the HA sites are protonated and only strong sites are able to bind cations. At basic pH, all sites are deprotonated and able to bind cations. Fig. 1b displays the proportion of Fe(II) bound to HA in the pH range 1.95 to 9.90 ($\text{DOC} = 48.5 \text{ mg L}^{-1}$). At $\text{pH} = 1.95$, only 4.1% of the total Fe(II) was complexed with HA. The adsorbed Fe(II) percentage then increased progressively with increasing pH to reach 51.3% at $\text{pH} = 4.99$ and $> 99\%$ at $\text{pH} > 8.13$. Fig. 1c displays the adsorbed Fe(II) percentage at $\text{DOC} = 76.5 \text{ mg L}^{-1}$ in the pH range 2.95 to 8.89. 17.6% of the total Fe(II) was bound to HA at $\text{pH} = 2.95$, with this amount reaching 62.9% and $> 99\%$ for pH values = 4 and 8.89, respectively. Both pH sorption edge experiments exhibited the same trend, but the percentages increased with the increasing DOC concentration.

3.2. Model results

As explained in section 2.4, we used the coupling PHREEPLOT-PHREEQC-Model VI to simultaneously fit all our datasets. We determined the following values for the Fe(II) binding parameters (without the linear relationship between $\log K_{\text{MA}}$ and $\log K_{\text{MB}}$): $\log K_{\text{MA}} = 2.19 \pm 0.16$, $\log K_{\text{MB}} = 4.46 \pm 0.47$ and $\Delta \text{LK}_2 = 3.90 \pm 1.31$ with $R^2 = 0.97$.

3.2.1. Adsorption isotherm

The root-mean-square error (RMSE) was calculated as $\text{RMSE} = \sqrt{\text{mean}(\log v_{\text{exp}} - \log v_{\text{calc}})^2}$ where v_{exp} and v_{calc} are the amounts of Fe bound to HA per gram of DOC for the experimental and modeled data, respectively. The modeled data

were plotted as a solid line in Fig. 1a. The fit goodness of the adsorption isotherm was validated by the RMSE value of 0.09. $\log [\text{Fe(II)}\text{-AH}]$ and $\log [\text{Fe(II)}]_{\text{UF}}$ increased as the $[\text{Fe(II)}]$ total concentration increased (Fig. 1a). The modeled data were close to the experimental data as shown by the RMSE equal to 0.08 for this dataset. In Fig. 1a, the first points, $\log [\text{Fe(II)}\text{-HA}] = -4.95$ and -5 , were both far from the modeled data compared to the other points. Despite the fact that the fit appears to be visually poor at low concentrations, the RMSE is good. This can be explained by the scale used here ($\log[\text{Fe(II)}]$), which reinforces the differences. We tried to improve the fit for the points at low concentrations using the new parameters, but the fits for the global pH adsorption edge were worse. Furthermore, these parameters did not involve any modifications of the Fe(II) speciation on the HA sites. Therefore, we chose the set of binding parameters that provided the best RMSE.

3.2.2. pH adsorption edge

The percentage of adsorbed Fe(II) increased with increasing pH in both experiments (DOC concentration = 48.8 mg L^{-1} , Fig. 1b and 76.5 mg L^{-1} , Fig. 1c). The RMSE for the first dataset (DOC = 48.8 mg L^{-1} , Fig. 1b) is equal to 0.11. The experimental percentage of adsorbed Fe(II) was higher than in the simulation at $\text{pH} = 6.94$ and 8.13 , but lower than in the simulation at $\text{pH} = 5.90$ and 2.99 , explaining this high RMSE. For the second dataset (DOC = 76.5 mg L^{-1} , Fig. 1c) the RMSE is much lower (0.06) indicating a better fit between the modeled and experimental data. In both cases, the pH adsorption edge showed low Fe(II) complexation by HA ($< 20\%$) at acidic conditions ($\text{pH} \leq 3$), the complexation becoming instead nearly quantitative ($> 90\%$) at $\text{pH} > 7$.

3.3. *Fe(II) speciation onto HA binding sites*

Speciation was calculated from the fit of the experimental data using PHREEQC-Model VI. PHREEQC-Model VI is able to calculate the proportions of Fe(II) bound to each site defined in the model: phenolic (sum of Fe(II) bound to phenolic monodentate, pheno-

phenolic bidentate and pheno-pheno-phenolic tridentate groups), carboxylic (sum of Fe(II) bound to carboxylic monodentate, carboxy-carboxylic bidentate and carboxy-carboxy-carboxylic tridentate groups), carboxy-phenolic (sum of Fe(II) bound to carboxy-phenolic bidentate, carboxy-carboxy-phenolic and carboxy-pheno-phenolic tridentate groups), etc. The complex denticity was also determined: monodentates correspond to the sum of Fe(II) bound to carboxylic and phenolic groups, bidentates to the sum of Fe(II) bound to carboxy-phenolic, pheno-phenolic and carboxy-carboxylic groups and tridentates to the sum of Fe(II) bound to carboxy-carboxy-carboxylic, carboxy-pheno-phenolic and pheno-pheno-phenolic groups.

3.3.1. Adsorption isotherm

The majority of Fe(II) (between 73.1 and 83.9% of Fe(II)) was bound to HA by bidentates. The remaining Fe(II) was mainly bound through tridentate sites (20%), as the amount of Fe(II) bound through tridentate sites is insignificant (< 1%, Fig. 2a). Moreover, the proportions of Fe(II) bound to carboxylic, carboxy-phenolic and phenolic functional groups were 50%, 40% and 10%, respectively (Fig. 2b). All Fe(II) bound to HA was in the form of Fe^{2+} (Fig. 2c).

3.3.2. pH adsorption edge

In the experiment conducted with a DOC concentration of 48.8 mg L^{-1} , more than 80% of Fe(II) bound to HA formed bidentates and approximately 15% formed tridentates. The number of monodentate sites was insignificant in this experiment (< 1%, Fig. 3a). The pH did not influence their distribution, but strongly controlled the nature of the functional groups involved. Thus, the proportion of carboxylic groups decreased from > 80% at pH = 1.95 to < 10% at pH = 9.90 (Fig. 3b), which was compensated by an increased role of carboxy-phenolic sites, which accounted for between 20 and 60% of the Fe(II) binding with increasing pH. Some phenolic sites were also involved (up to 20% at pH = 9.90), but only at high pH. From acidic to neutral pH conditions, all Fe(II) bound to HA occurred as Fe^{2+} (Fig. 3c). The Fe-OH^+ species appeared only from pH = 8.13 (1.35% of Fe(II) bound to HA), with

the proportions of this species then increasing with increasing pH to reach 38.5% at pH = 9.90.

In the experiment conducted with a DOC concentration of 76.5 mg L⁻¹, considering the nature and proportion of the functional groups involved, the evolution with regards to the pH was quite similar to that observed in the previous experiment. Approximately 80% (from 78.8 to 86.7%) of Fe(II) was bound to HA as bidentates (Fig. 4a) and approximately 20% as tridentates. The proportions of Fe(II) bound as monodentates were insignificant (< 1%). At acidic pH (2.95 and 4.95), 66% of Fe(II) was bound to carboxylic groups, approximately 30% to carboxy-phenolic groups and approximately 1% to phenolic groups (Fig. 4b). At pH = 6.93, 52.3% of Fe(II) was bound to carboxy-phenolic groups, 33.5% to carboxylic groups and 14.2% to phenolic groups. At basic pH (8.89), Fe(II) was more bound to carboxy-phenolic groups (59.6%), than to phenolic (21.8%) or carboxylic groups (18.6%). At acidic and neutral pH (pH 2.95, 4.95 and 6.93), Fe(II) was completely bound to HA as Fe²⁺ (Fig. 4c), whereas at basic pH (8.89), 6% of Fe(II) bound to HA occurred as Fe(OH)⁺.

4. Discussion

4.1. Validation of the set of binding parameters

Two strategies were used to validate the new estimated binding parameters (log K_{MA} and log K_{MB}). They were first compared with the parameters estimated by the LFER technique. Second, the conditions of the previously published experimental studies were put into PHREEQC-Model VI to calculate the expected Fe(II)-HS proportions; the calculated proportions were then compared with the proportions determined experimentally.

The binding parameters determined in the present work are different from those determined by Tipping (1998) due to (1) the deletion of the linear relationship between log K_{MA} and log K_{MB} as performed by Marsac et al. (2011) and (2) the experimental dataset used for fitting. Tipping (1998) only used the criticized Van Dijk (1971) dataset. Van Dijk (1971) studied Fe(II) binding with HA through titrations and consequently only provided pH data.

The specific $\log K_{MA}$ and $\log K_{MB}$ parameters can be estimated using the linear relationship existing between these parameters and the acid acetic ($\log K_{MA}$), first hydrolysis ($\log K_{MA}$), and catechol ($\log K_{MB}$) stability constants. The stability constant of Fe(II) with acetic acid ($\log K(\text{Fe(II)-AA})$) and the first hydrolysis constant of Fe^{2+} ($\log K(\text{Fe(II)-OH})$) are $\log K(\text{Fe(II)-AA}) = 3.32$ and $\log K(\text{Fe(II)-OH}) = 7.72$ (IUPAC). Using these values, the LFERs determined by Pourret et al. (2007) and Marsac et al. (2011) resulted in the following estimates for the $\log K_{MA}$ and $\log K_{MB}$ values: $1.57 < \log K_{MA} < 2.35$ (acetic acid LFER, $R^2 = 0.80$) and $1.69 < \log K_{MA} < 2.65$ (first hydrolysis LFER, $R^2 = 0.78$); $3.70 < \log K_{MB} < 4.09$ (catechol LFER, $R^2 = 0.95$). As can be seen, there is a full agreement between the $\log K_{MA}$ value determined experimentally in this study (2.19) and the $\log K_{MA}$ values estimated using the LFER method. For $\log K_{MB}$, the experimentally determined value (4.46) is bigger than the LFER estimate. However, the shift remains limited, as it was only 15%.

An attempt to experimentally determine the binding constants of Fe(II) by FA was made by Schnitzer and Skinner (1966) using an ion-exchange method. Two sets of experiments were performed with the same concentration of Fe(II) ($1.79 \cdot 10^{-3} \text{ mol L}^{-1}$): one at pH 3.5 with 201, 402, 603, 804 and 1005 mg L^{-1} of FA, and one at pH = 5 with 100.5, 120.6, 160.8, 201 and 241.2 mg L^{-1} of FA (Fig. 5). They showed that Fe(II) binding with FA was only slightly affected by pH. Schnitzer and Skinner's (1966) dataset was calculated using our new Fe(II)-HA binding parameters with PHREEQC-Model VI. For this purpose, we used the relationship between $\log K_{MA}(\text{HA})$ and $\log K_{MA}(\text{FA})$ established by Tipping (1998): $\log K_{MA}(\text{HA}) = 0.64 * \log K_{MA}(\text{FA}) + 0.79$ ($R^2 = 0.84$). We also used this relationship to determine the binding parameters for FA: $\log K_{MA} = 2.19$; $\log K_{MB} = 4.46$ and $\Delta \text{LK}_2 = 3.89$. At pH = 3.5, the modeled partitioning of Fe(II) between free species and FA complexes was close to the experimental one, with a low RMSE value of 0.04, but the fit was not as good at pH = 5 as the RMSE value was much higher (RMSE = 0.26) (Fig. 5). Thus, the new set of binding parameters failed to reproduce Schnitzer and Skinner's data (1966) over the entire pH range. It is to be noticed, however, that Tipping (1998) himself did neither use Schnitzer and Skinner's (1966) dataset in his estimation of Fe(II)-FA binding parameters, nor for the other

metals studied by these authors. Tipping and Hurley (1992) justified this by noting that the $\log K_{MA}$ value of Fe(II)-FA calculated from Schnitzer and Skinner's (1966) dataset did not rank properly when compared to the $\log K_{MA}$ values determined for the other cations. The Fe(II) $\log K_{MA}$ obtained using Schnitzer and Skinner's (1966) dataset was intermediate between Pb^{2+} and Cu^{2+} $\log K_{MA}$, in contradiction with the findings of Martell and Smith (1977), who established that Fe^{2+} stability constants should be lower than those for Pb^{2+} and Cu^{2+} binding constants with mono- and di-carboxylic acids. This dataset was also criticized by Milne et al. (2003) who attributed it a poor score when trying to fit it using the NICA-Donnan model. They could not simply extrapolate the calculated data from Schnitzer and Skinner's (1966) experimental data and had to constrain the second site distribution values using LFER between their binding parameters and $\log KOH$. According to Tipping and Hurley (1992) it is possible that in the experimental conditions used by Schnitzer and Skinner (1966), part of the Fe(II) was oxidized into Fe(III), and that this oxidation caused the observed shift between the calculated and expected $\log K_{MA}$ values.

Yamamoto et al. (2010) estimated the binding capacities of HS with Fe(II) using a colorimetric method. Yamamoto et al. (2010) used HA and FA extracted from compost with a HA concentration of 1 mg L^{-1} and determined the Fe(II) concentrations using the ferrozine colorimetric method. As with Schnitzer and Skinner's (1966) data, the new binding parameters obtained here failed to model these data ($RMSE > 1$). This failure is thought to be due to an analytical bias in the dataset, related to the colorimetric method. Jackson et al. (2012) emphasized the poor reliability of the colorimetric methods to determine the Fe(II) concentration bound to HS.

Rose and Waite (2003) studied the kinetics of Fe(II) complexation with NOMs. They used 12 different NOMs recovered from the soil leaf litter layer and one fulvic acid stock solution: Suwannee River FA (SRFA). Binding experiments were performed in seawater with a nominal salinity of 36 (ionic strength $\approx 1.7 \text{ M}$) at pH 8.1. Three Fe(II) concentrations were used: 0.25, 0.5 and $1 \text{ } \mu\text{mol L}^{-1}$ with a total organic carbon concentration of 1 mg L^{-1} . All of the data used for the calculations were equilibrium concentrations (i.e. concentrations

determined at the end of the kinetics). The amount of Fe(II) adsorbed at equilibrium differed from one NOM to another. For example, for $[\text{Fe(II)}]_{\text{tot}} = 1 \mu\text{mol L}^{-1}$, the percentages of experimentally adsorbed Fe(II) varied from 5 to 100% depending on the NOM sample considered. This variability could not be reproduced by PHREEQC-Model VI (Fig. 6a), suggesting that some important properties of the NOM samples used were not taken into account by the model. In fact, the NOM samples used by Rose and Waite (2003) in their experiments were extracted from soil leaf litter, which were probably different in composition from the HS structure considered in PHREEC-Model VI. The situation is obviously much different for the SRFA sample used by Rose and Waite (2003), which has a structure that is typical for a HS. Interestingly, a reasonably good fit (RMSE = 0.03) was obtained with the new binding parameters for the data (SRFA) published by Rose and Waite (2003) (Fig. 6b). Note that data from Van Dijk's dataset (1971) were not used for the purpose of a validation test considering, as Milne et al. (2003) did, that the measurement of the pH values was not sufficiently precise in this study.

Thus the new binding parameters obtained here were able to model the Fe(II) adsorption onto both HA and FA reasonably well. The new set of parameters satisfactorily reproduced Rose and Waite's (2003) experimental data for SRFA (RMSE = 0.03) and the present data. It is interesting to see that previously determined binding parameters used a linear relationship between $\log K_{\text{MA}}$ and $\log K_{\text{MB}}$ (Tipping, 1998). In fact, the present data cannot be modeled if this relationship is imposed. As Marsac et al. (2012) did, we had to remove this constraint to satisfactorily model our data. When Marsac et al. (2012) tried to model their data using the linear relationship, REE patterns were identical for weak and strong sites, which is impossible with regards to their stability constants with single organic ligands, such as acetate (weak sites) or EDTA (strong sites) (Marsac et al., 2011). To reproduce correctly REE patterns binding to strong sites, the authors had to delete this linear relationship. Consequently, $\log K_{\text{MB}}$ values decreased and were compensated by higher values of $\Delta\text{LK}_{2\text{C}}$ and $\Delta\text{LK}_{2\text{P}}$. Most likely, this explains why we also obtained a high ΔLK_2 value (3.90). Despite the use of a glovebox, part of the Fe(II) might be oxidized into Fe(III) which

might explain the strong Fe adsorption at acidic pH and the subsequent high value of ΔLK_2 . However, humic substances are expected to delay the oxidation of Fe (Wolthoorn et al., 2004; Kleber et al., 2005; Weber et al., 2006; Miller et al., 2009) and no Fe(III) was measured in the ultrafiltrated solution. Indeed, Weber et al. (2006) performed sorption experiments of Fe(III) (50 and 30 μM) with organic matter (1.8 g L⁻¹). They showed that for the lowest total Fe concentrations, a fraction of Fe can be reduced by humic substances. The concentrations of Fe(II) used in the present study were from 11 to 56 μM , within the range of Weber et al.'s (2006) experiments. Weber et al.'s (2006) study suggests therefore that OM tends to reduce Fe(III) into Fe(II) and not to oxidize it. So, oxidation of Fe(II) into Fe(III) seems unlikely in the experimental conditions used in the present study.

However at any rate, the high ΔLK_2 value obtained here is lower than the ΔLK_2 of 4.0 reported by Marsac et al. (2012; 2013) for the binding of Fe(III) by HS (Table 2), which is rather consistent given that Fe(III) species are known to form stronger complexes with HS than Fe(II) species (Ou et al., 2009). The new binding parameters determined in the present study ($\log K_{MA} = 2.19$, $\log K_{MB} = 4.46$ and $\Delta LK_2 = 3.90$) are therefore in accordance with the Fe(III) binding parameters of Marsac et al. (2013).

4.2. Fe(II) speciation onto HA binding sites

Although several extended X-ray absorption fine structure (EXAFS) analyses concerning the binding of Fe to OM have been performed for Fe(III) (Gustafsson et al., 2007; Van Schaik et al., 2008; Morris et al., 2012), no spectroscopic data are currently available for Fe(II)-OM binding. Therefore, the validation of the speciation results obtained here for Fe(II)-HS binding using PHREEQC-Model VI can only be achieved through comparison with the case of Fe(III). Gustafsson et al. (2007) analyzed the binding of Fe(III) to organic soils using EXAFS spectroscopy. They determined that Fe(III) bound to organic soils seems to occur through dimeric and trimeric complexes, where Fe ions are connected by μ -oxo bridges. Using scanning electron microscopy (SEM) images, Ou et al. (2009) suggested that Fe(III) is

bound to HA as oligomeric species (not as single ions). These results differ from the EXAFS studies performed on the binding of Fe(III) with FA (Van Schaik et al., 2008), peat humic acid (Karlsson et al., 2010) and organic soils (Karlsson et al., 2010), which have shown the formation of mononuclear complexes between Fe(III) and organic functional groups. This reveals that mononuclear Fe species are likely the dominant species involved in the binding of Fe(III) by HS. The PHREEQC-Model VI binding hypothesis considers that Fe(II) is bound to HS as mononuclear species, which appears consistent with the spectroscopic data obtained for Fe(III). Moreover, Gustafsson et al. (2007), Van Schaik et al. (2008) and Morris and Hesterberg (2012) suggested that the functional groups involved in the binding of Fe(III) are phenolate and carboxylate groups, while Weber et al. (2006) argued for a dominant role of carboxylic groups. They tested two models for Fe(III)-HA binding with or without a contribution from the phenolic groups. With phenolic groups, the adsorbed Fe(III) with phenolate was negligible. Using Fourier transform infrared spectroscopy (FTIR), Ou et al. (2009) suggested that Fe(III) is mainly bound to HA carboxylate and carboxylic groups via monodentate and bidentate sites, whereas binding with HA phenolate and carboxy-phenolate groups would occur mainly through monodentate and bidentate sites, respectively. The results of the modeling calculations performed in this study indicated that Fe(II) could bind to HA by carboxylic and phenolic groups as suggested by Gustafsson et al. (2007), Van Schaik et al. (2008) and Morris and Hesterberg (2012) for Fe(III). Moreover, modeling calculations revealed that Fe(II) was bound with HA mainly through bidentate complexes with a subordinate participation of tridentate sites. At basic to neutral pH, Fe(II) was bound to carboxylic groups, which is consistent with the results obtained for Fe(III) by Weber et al. (2006) and Ou et al. (2009). By contrast, as suggested by Ou et al. (2009) for Fe(III), Fe(II) was mainly bound to carboxy-phenolic groups.

4.3. Environmental implications

These new data provided the assessment that Fe(II) could be strongly bound to HA. Iron(II) complexation with HA or OM therefore put strong constraints on the observed dissolved Fe(II) concentrations and the possible transfer of Fe(II) within waters and hydrosystems enriched in OM. For example, in a coastal environment, Fe occurs mainly as insoluble Fe-oxyhydroxides (Rose and Waite, 2003). Bioavailable Fe is therefore relatively scarce in such environments. However, any Fe(II) binding by OM can largely increase its solubility by preventing its hydrolysis/oxidation, therefore forming mobile soluble organic complexes (Rose and Waite, 2003; Miller et al., 2009; 2012). These studies showed that Fe(II)-OM binding forms strong complexes, which can have a significant effect on its solubility in coastal waters.

Another major environmental implication concerns the behavior of Fe(II) within wetlands. When reductive conditions develop in wetlands during the flooding season, high amounts of dissolved Fe(II), DOC and trace elements can be released in the soil solution (Ponnamperuma, 1972; McBride, 1994; Dia et al., 2000; Oliv  -Lauquet et al., 2001). In such conditions, with regards to the strong affinity of Fe(II) for OM, Fe(II) will be mainly bound to DOC thus preventing its precipitation as a secondary mineral such as magnetite or green rust. As the formation of Fe-rich secondary minerals is limited, Fe-associated trace elements are not taken up and thereby remain highly mobile. Such processes have been previously suggested by Davranche et al. (2013) from kinetic modeling. These authors showed that organic-mediated Fe(II) complexation is a major controlling factor of Fe reactivity in wetland soils.

Moreover, the presence of other cations can change the quantities of Fe(II) bound to HA. The values of the binding parameters for some cations (e.g. Mg(II), Ca(II), Mn(II), Co(II), Ni(II), Zn(II), Sr(II), Cd(II), Ba(II)) are smaller than those of Fe(II), which means that the presence of these cations would only have minor consequences on the quantities of Fe(II) bound to HA or FA. By contrast, some other cations (Al(III), Cr(III), Fe(III), Pb(II), REE) have

binding parameters values that are higher than those of Fe(II) and can therefore strongly compete with Fe(II) for HA binding if they are present in sufficient concentrations.

Finally, several authors have shown that Fe(III), through the formation of ternary complexes with HS, allows the binding of As(V) as oxyanion to OM (Sharma et al., 2010) and HS (Mikutta et al., 2011). In anoxic environments, as Fe(II) is the dominant species, such ternary associations are thus expected to be formed binding Fe(II), HS and As(III) or As(V). In this context, the present study and the new set of binding parameters suggested that such ternary associations with Fe(II) may be relatively important in wetlands and floodplains. Thus these data constitute a first step for studying the formation of a cationic bridge between As(III, V), Fe(II) and HA and therefore the involved processes controlling As mobility at the soil-water interface within organic-rich environments undergoing redox alternations.

5. Conclusions

Iron(II) adsorption experiments onto humic acid (HA) were designed to study the binding of Fe(II) by organic matter. The experimental results showed that Fe(II) was strongly bound to HA as previously observed by several authors (Schnitzer et al., 1966; Van Dijk, 1971; Tipping, 1998; Rose and Waite, 2003; Yamamoto et al., 2010). Coupling PHREEPLOT and PHREEQC-Model VI, a new set of specific binding parameters was calculated for Fe(II)-HA complex ($\log K_{MA} = 2.19 \pm 0.16$, $\log K_{MB} = 4.46 \pm 0.47$ and $\Delta LK_2 = 3.90 \pm 1.31$). These binding parameters are higher than those previously calculated by Tipping (1998) using only Van Dijk's dataset (1971). The new set of Fe(II)-HA binding parameters was used to test and validate the model using the relationship between the HA and FA parameters. They reasonably reproduced Rose and Waite's data (2003) on the adsorption kinetics of Fe(II) with SRFA. The RMSE between the calculated and experimental data was low (0.03), validating the newly determined set of binding parameters as well as the experimental and modeling approaches used in this study. This new dataset of binding parameters implies a higher complexation between Fe(II) and humic substances than that previously suggested, and

shows that OM is of major importance in the fate and bioavailability of Fe(II) and possibly associated trace metals such as As within the environment.

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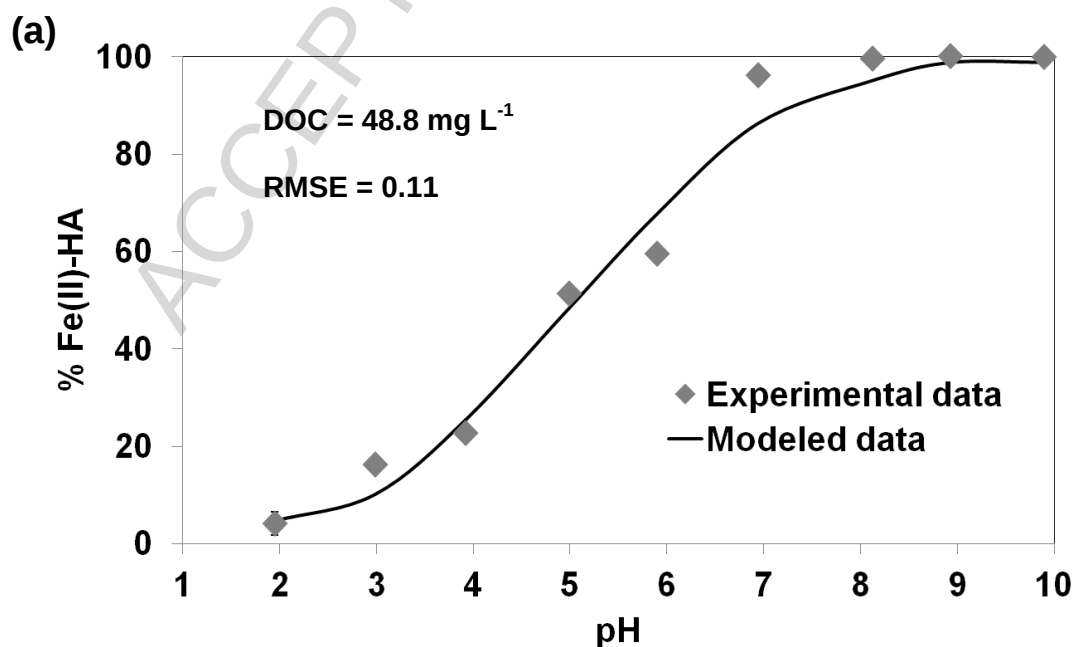
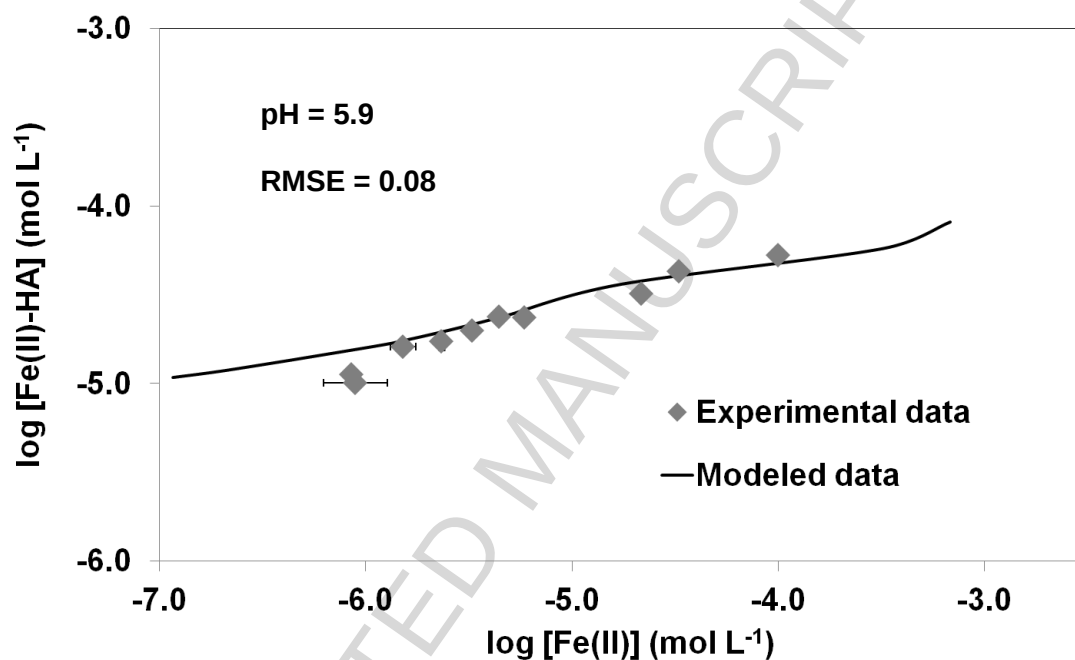
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Figures:

As each experiment was performed in triplicate, each point represents the average of the triplicates. The error bars represent the standard deviation (in log or percentage, depending on the data) calculated from the triplicated experimental data.



(b)

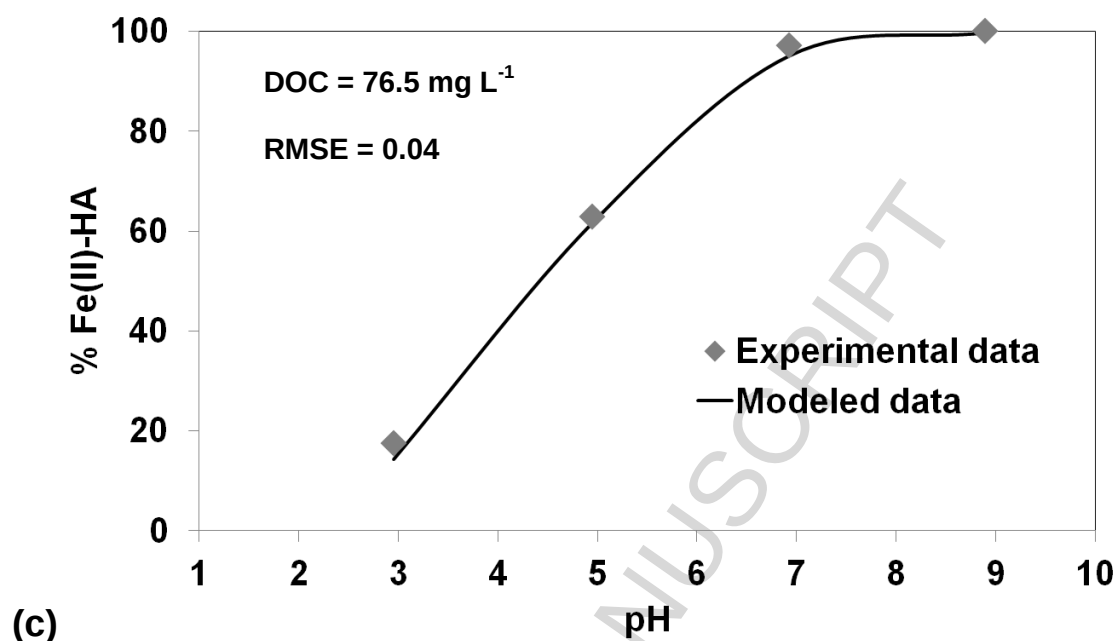
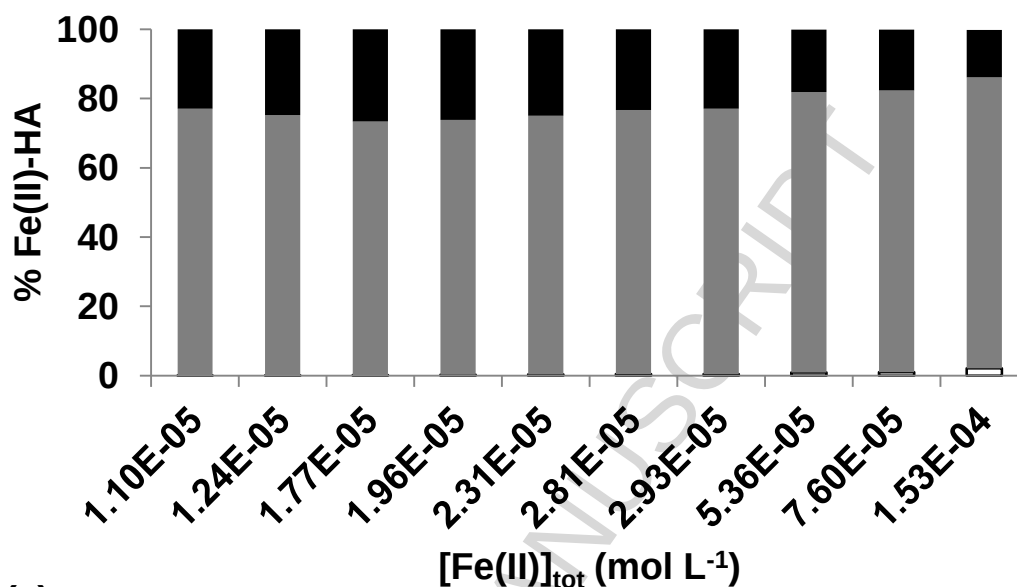
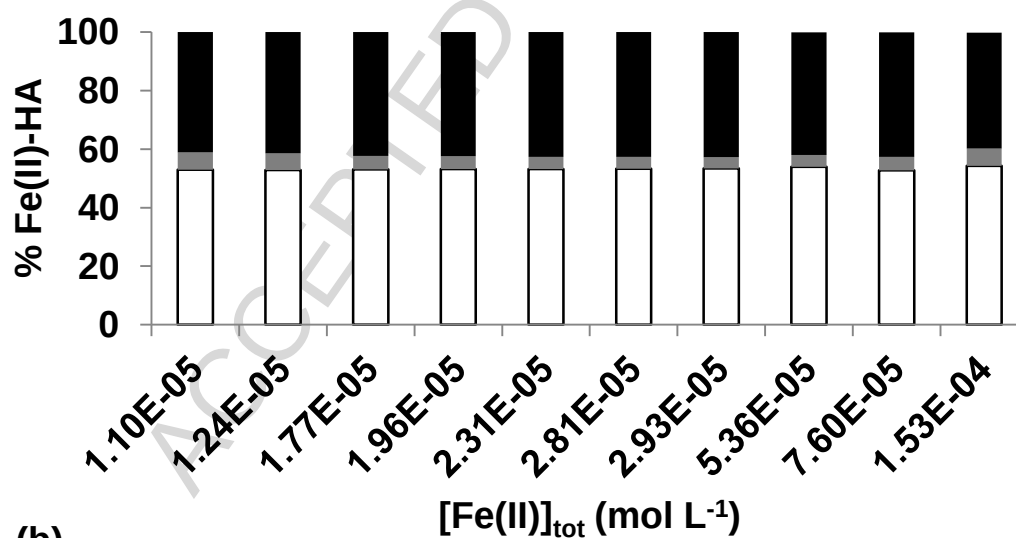


Fig. 1: (a) Adsorption isotherm: experimental and modeled data, (b) pH adsorption edge - DOC = 50 mg L⁻¹ (c) pH adsorption edge - DOC = 76 mg L⁻¹ (the error bars calculated from the triplicated experiments are within the symbols.).



(a)

□ Monodentates ■ Bidentates ■ Tridentates



(b)

□ Carboxylic sites ■ Phenolic sites ■ Carboxy-phenolic sites

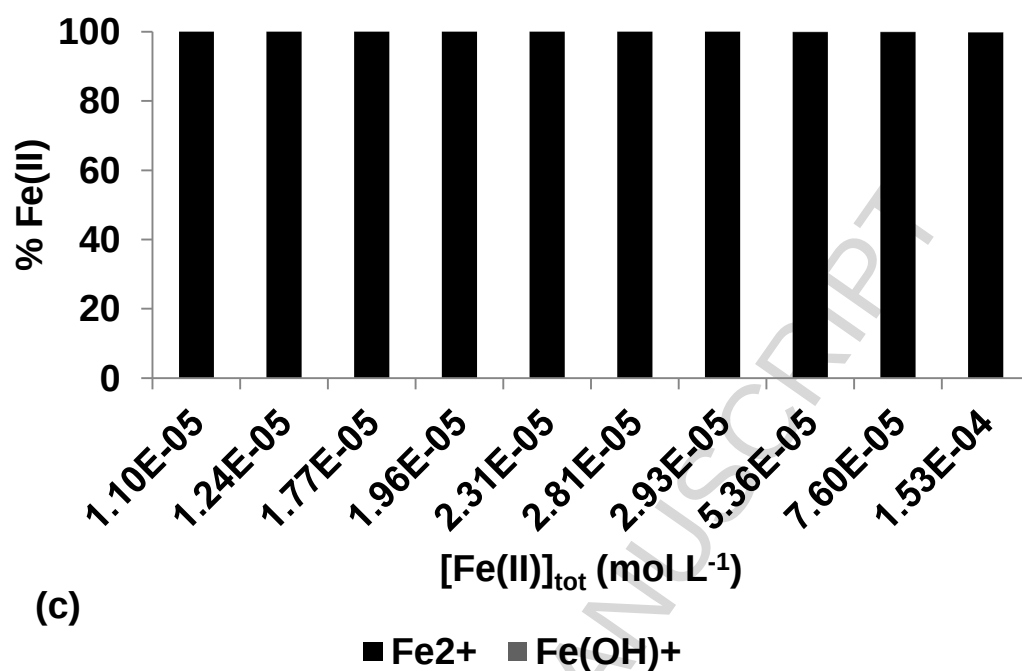
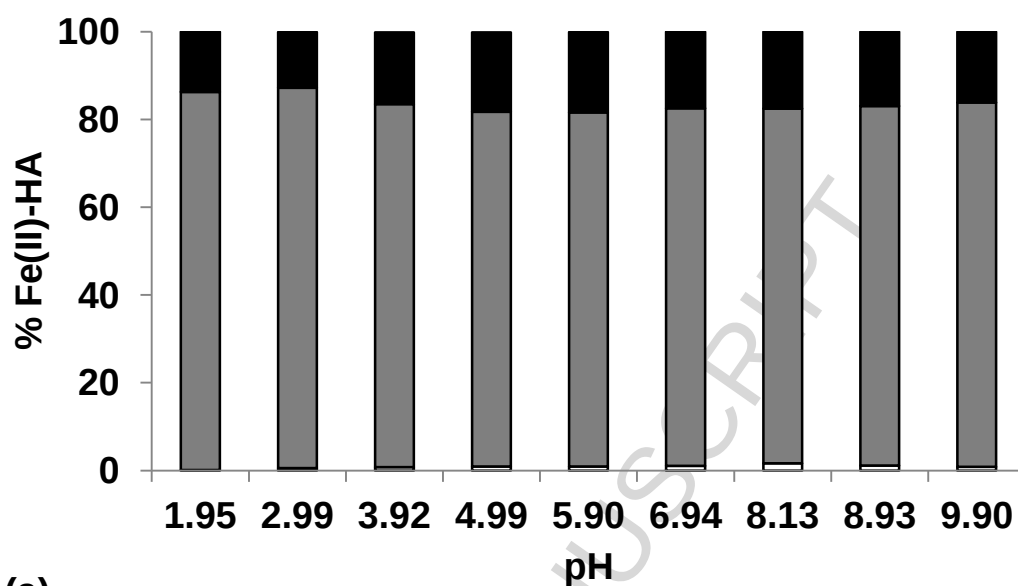
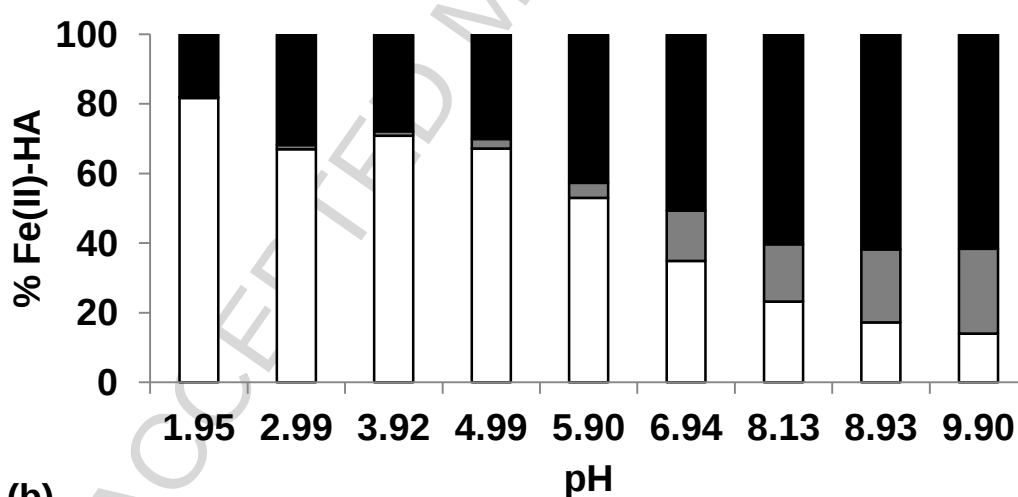


Fig. 2 Fe(II) speciation onto the binding sites - adsorption isotherm (a) Denticity (b) Type of site (c) Form of adsorbed Fe(II).



(a)

□ Monodentates ■ Bidentates ■ Tridentates



(b)

□ Carboxylic sites ■ Phenolic sites ■ Carboxy-phenolic sites

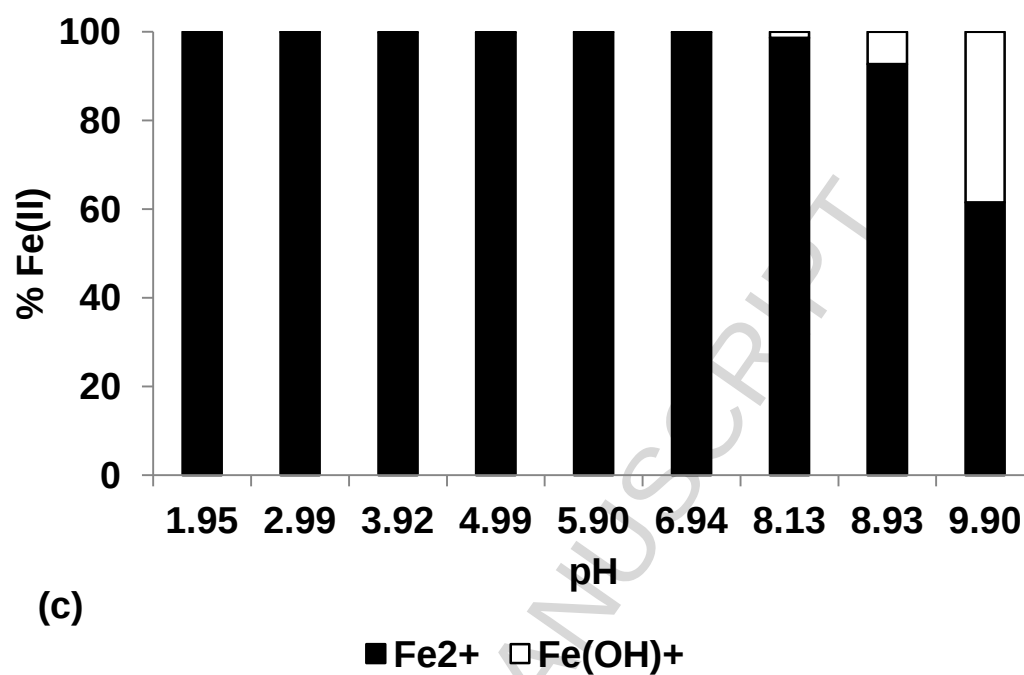
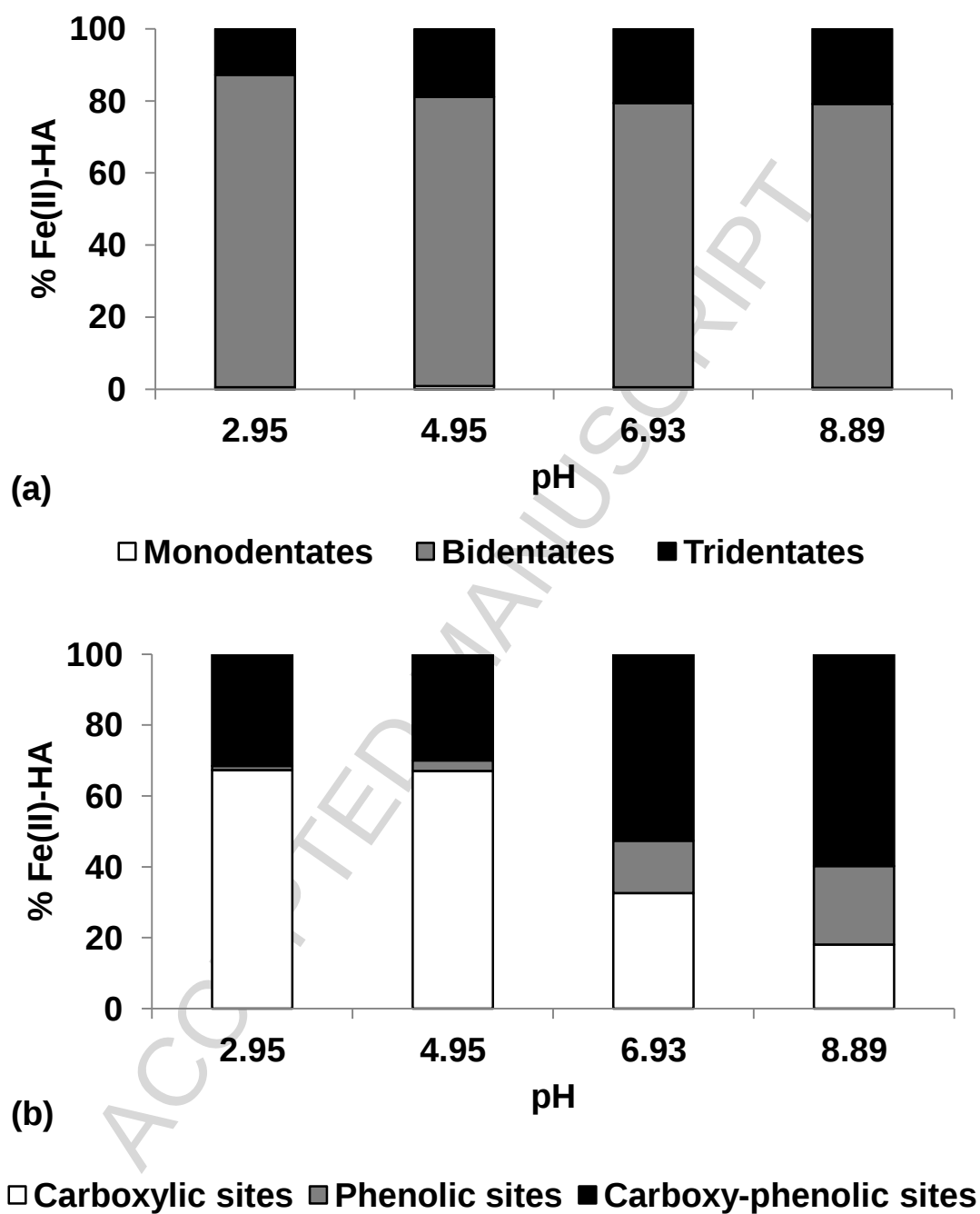


Fig. 3 Fe(II) speciation onto the binding sites - pH adsorption edge - DOC = 50 mg L⁻¹ (a) Denticity, (b) Type of site, (c) Form of adsorbed Fe(II).



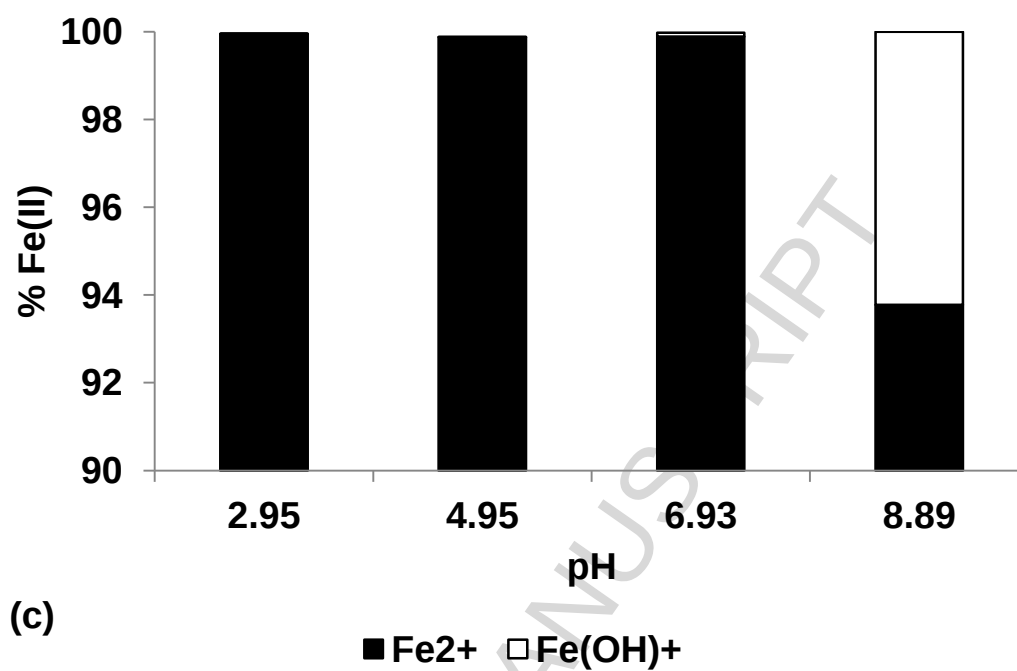


Fig. 4 Fe(II) speciation onto the binding sites - pH adsorption edge - DOC = 76 mg L⁻¹ (a) Denticity, (b) Type of site, (c) Form of adsorbed Fe(II).

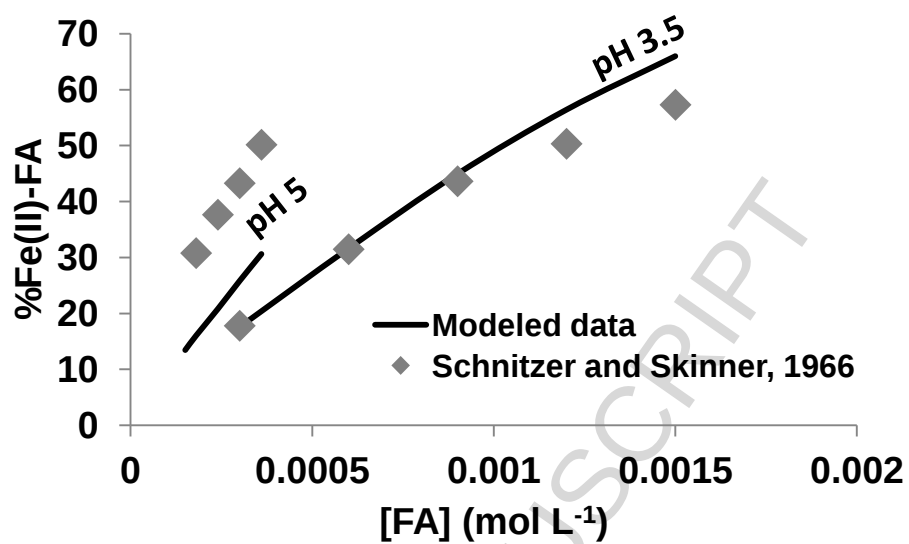


Fig. 5: Schnitzer and Skinner's dataset: experimental and modeled % of Fe(II) adsorbed onto FA.

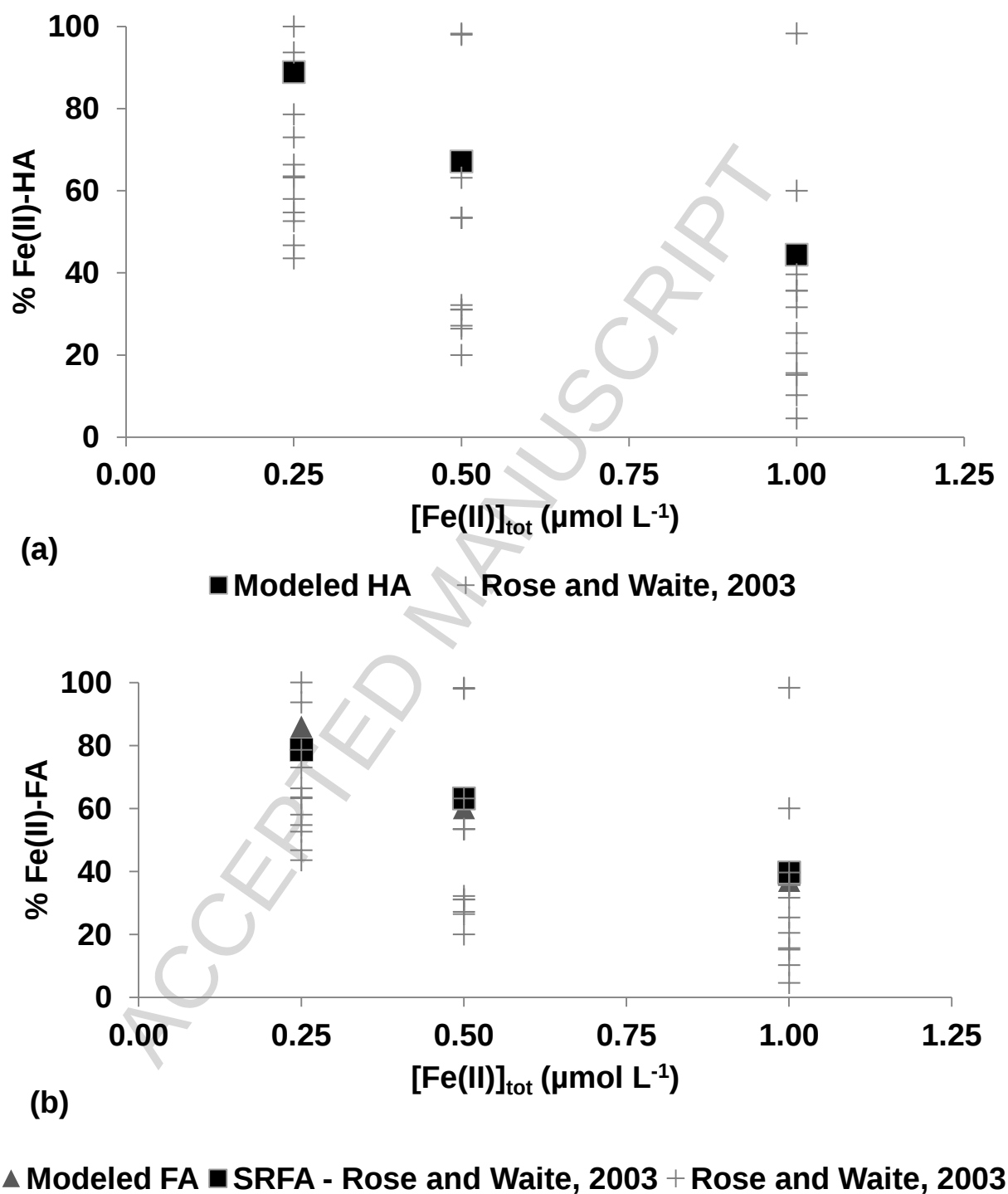


Fig. 6 Rose and Waite's dataset: (a) Comparison % of Fe(II) adsorbed onto twelve different natural organic matters through an experimental approach and HA modeled data, (b) Experimental (SRFA)-mediated and modeled % of Fe(II) adsorbed onto FA.

Tables

Table 1

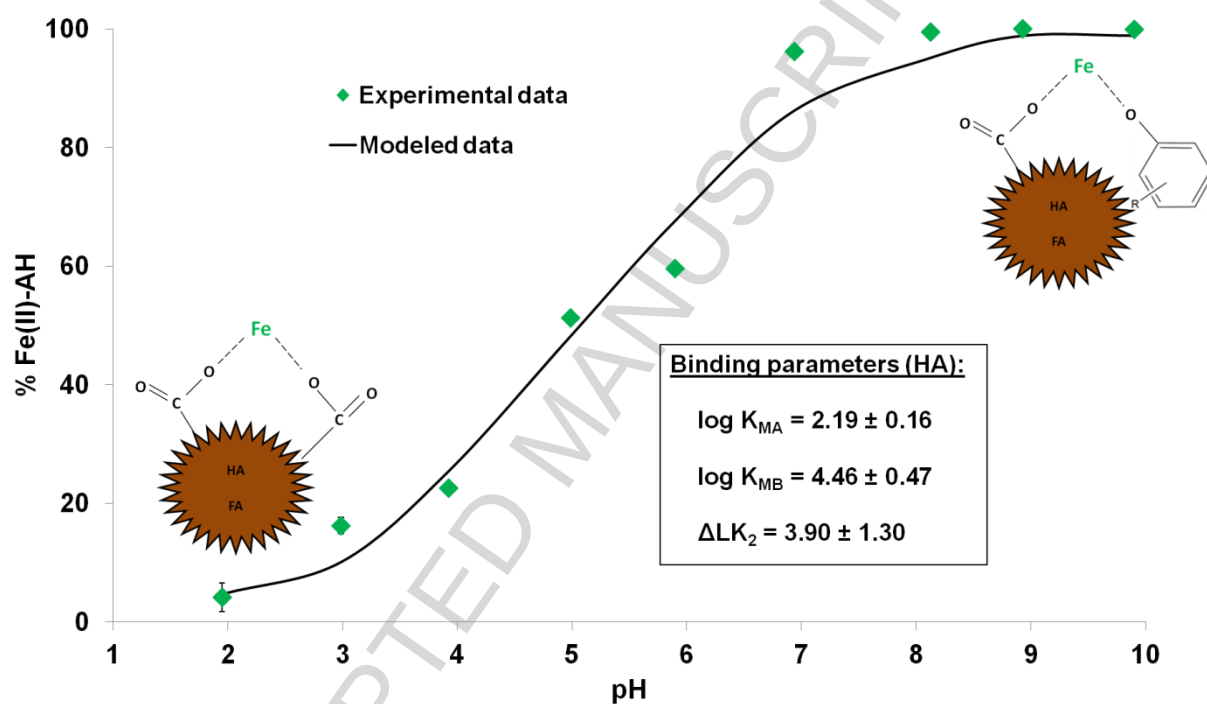
PHREEQC-Model VI parameters for the humic and fulvic acids (Tipping, 2002)

Parameter	Description	Values	
		HA	FA
n_A	Abundance of type A sites (mol g^{-1})	0.0033	0.0048
n_B	Abundance of type B sites (mol g^{-1})	0.00165	0.0024
pK_A	Intrinsic proton dissociation constant for type A sites	4.1	3.2
pK_B	Intrinsic proton dissociation constant for type B sites	8.8	9.4
ΔpK_A	Distribution term that modifies pK_A	2.1	3.3
ΔpK_B	Distribution term that modifies pK_B	3.6	4.9
f_B	Fraction of proton sites that can make bidentate sites	0.5	0.42
f_T	Fraction of proton sites that can make tridentate sites	0.065	0.03
$\log K_{MA}$	Intrinsic equilibrium constant for metal binding at type A sites	Fitted from experimental data	
$\log K_{MB}$	Intrinsic equilibrium constant for metal binding at type B sites	Fitted from experimental data	
ΔLK_{1A}	Distribution term that modifies $\log K_{MA}$	-0.7	0.5
ΔLK_{1B}	Distribution term that modifies $\log K_{MB}$	0.8	2.1
ΔLK_2	Distribution term that modifies the strength of bidentate and tridentate sites	Fitted from experimental data	

Table 2: $\log K_{MA}$, $\log K_{MB}$, ΔLK_{2C} and ΔLK_{2P} values of the different species of Fe from Tipping (1998) (1) from this study (2), and from Marsac et al., 2013 (3).

Parameter	$\text{Fe}^{2+} / \text{FeOH}^+ (1)$	$\text{Fe}^{2+} / \text{Fe(OH)}^+ (2)$	$\text{Fe}^{3+} / \text{FeOH}^{2+} (1)$	$\text{Fe}^{3+} / \text{FeOH}^{2+} (3)$
$\log K_{MA}$	1.3	2.19	2.5	3.5
$\log K_{MB}$	3.257	4.46	7.325	6.9
ΔLK_{2C}	0.81	3.9	2.2	4/0
ΔLK_{2P}	0.81	3.9	2.2	4/0

Graphical Abstract:



Highlights:

- We measured Fe(II)-Humic acid adsorption isotherm and pH sorption edge
- Results were modeled using a coupling of PHREEPLOT-PHREEQC-Model VI
- We determined the binding parameters for Fe(II)-humic acid complex
- We validated the binding parameters using LFERs and published datasets
- **We determined the speciation of Fe(II)-humic acid complexes using PHREEQC-Model VI**