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Originally published:

April 2023

### Journal of Materials Research 38(2023), 2752-2763

DOI: https://doi.org/10.1557/s43578-023-00998-8

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#### Assessing the reactivity of Fe(II) sorbed on smectite clays: U(VI) reduction

Journal:	Environmental Science & Technology
Manuscript ID	es-2019-068495
Manuscript Type:	Article
Date Submitted by the Author:	12-Nov-2019
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2	clays: U(VI) reduction
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#### 22 ABSTRACT

23 The reactivity of Fe(II) sorbed on three Ca-exchanged smectite clays was probed via U(VI) 24 reduction at pH 6.0 ( $\pm$  0.2) under CO<sub>2</sub>-free, anoxic (O<sub>2</sub> <1 ppmv) condition. The clays varied 25 with regard to structural Fe content from Fe-free (0 wt.%) montmorillonite (MONT), to Fe-poor 26 (2.6 wt.%) montmorillonite (Fe-MONT), to Fe-rich (25.8 wt.%) nontronite (NAu-2). The U L<sub>III</sub>-27 edge XANES spectra showed no reduction of U(VI) in presence of Fe(II) sorbed on either Fe-28 MONT or NAu-2 after 72 h but a partial reduction (21%) on MONT after 24 h. U  $L_{III}$ -edge 29 EXAFS spectra further showed the formation of a soddyite-like surface precipitate on MONT in 30 presence of sorbed Fe(II) after 72 h. The Mössbauer data further reveals that 10 ( $\pm$  2) % of the 31 total sorbed Fe(II) was oxidized in presence of MONT before and an additional 6 ( $\pm$  2) % after 32 addition of U(VI). The mechanism of U(VI) reduction involves Fe(II) specifically sorbed on 33 oxidized and reduced strong edge sites of MONT. The non-reactivity of Fe(II) sorbed on Fe-34 MONT and NAu-2 towards U(VI) reduction is likely linked to the inter-valence charge transfer 35 (IVCT) between surface Fe(II) and structural Fe(III). The present study demonstrates that the 36 reduction capacity of Fe bound to the surface depends strongly on the nature of clay and 37 correspondingly on the oxidation state of the sorbed Fe. Thus Fe(II) sorbed clay cannot be 38 always considered as "universal reductant" for U(VI).

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40

41 BRIEF: Clay catalyzed redox processes between aqueous U(VI) and surface bound Fe(II).

#### 42 Introduction

The mobility and transport of hexavalent uranium, U(VI) in presence of clay have been studied for many years as possible remediation options in areas contaminated by uranium as well as for the evaluation of the performance of highly radioactive nuclear wastes (HLNW) repositories. As Fe(II) is common in both environments, the possible reduction of U(VI) to U(IV) and a comprehensive understanding of the Fe(II)-U(VI)-clay ternary system is necessary to mitigate the future possible risk in an anoxic uranium-polluted subsurface and to ensure long term stability of uranium at underground waste disposal sites.

50 Montmorillonite and nontronite are members of the smectite sub-group clay with structures 51 similar to 2:1 phyllosilicates and are known to sorb Fe(II) and U(VI) present in solution. The 52 mechanisms of U(VI) sorption on montmorillonite have been extensively investigated using 53 surface complexation modeling.<sup>1-6</sup> spectroscopic analysis and molecular dynamics simulation methods,<sup>7-14</sup> while the information available on the nontronite system is limited.<sup>15-17</sup> Surface-54 55 bound Fe(II) commonly behaves as stronger reductant than corresponding aqueous species in 56 both biotic and abiotic pathways, and therefore can transform organic (nitroaromatic explosives, 57 pesticides and polyhalogenated solvents) and inorganic (Cr, Tc, Se, U) pollutants into nontoxic forms.18-20 58

The sorption of Fe(II) on montmorillonite has been well documented.<sup>21-23</sup> The sorption affinity of pH-dependent montmorillonite makes it possible to distinguish three types of edge sites: weak sites, oxidized strong sites and reduced strong sites. Strong "oxidized" sites refer to those that are linked with heterogeneous surface oxidation of Fe(II), while on strong "reduced" and weak sites only sorption takes place.<sup>21,23</sup> The sorption of Fe(II) on iron-rich nontronite (NAu-1 & NAu-2) has been studied in detail by Tsarev et al.<sup>15</sup> and Jaisi et al.<sup>24</sup>. In addition to pH dependence, both 65 montmorillonite and nontronite undergo heterogeneous surface oxidation of sorbed Fe(II) via 66 different mechanisms. In montmorillonite, surface oxidation of Fe(II) may result in the formation of ferric hydroxide and lepidocrocite at lower and higher pH respectively<sup>25</sup> which, however, 67 coupled to catalyze the reduction of U(VI),<sup>15,25</sup> Se(IV),<sup>19</sup> 4-chloronitrobenzene.<sup>26,27</sup> In the case of 68 69 surface oxidation of Fe(II) on structural Fe(III)-bearing smectites, Fe(III)-montmorillonite and 70 nontronite have a unique characteristic known as inter-valence charge transfer (ICVT) between 71 sorbed Fe(II) and structural Fe(III). It has been proposed that Fe(III), which exists in clay 72 tetrahedral (nontronite) and/or octahedral (montmorillonite) sites participates in the electron 73 transfer reaction by a charge-shuttling process for redox mediation and catalysis or by edge and 74 basal sites. Therefore, various secondary mineral phases like green rust, ferrihydrite, lepidocrocite and magnetite may be formed as oxidation products.<sup>28-31</sup> The reactivity of sorbed 75 76 Fe(II) on structural Fe(III) bearing clays has been shown to vary in different cases. Jones et al.<sup>27</sup> 77 observed a significant decrease of reactivity of Fe(II) sorbed on nontronite after 18 days in reduction of 4-chloronitrobenzene. Tsarev et al.<sup>15</sup> observed a partial reduction of U(VI) by Fe(II) 78 79 sorbed on nontronite (NAu-1 and NAu-2) in presence of CO<sub>2</sub> while a partial oxidation of As(III) 80 to As(V) was reported by Ilgen et al.<sup>32</sup>

Our previous study<sup>25</sup> has addressed the pH dependent abiotic reduction of U(VI) by Caexchanged montmorillonite (Fe-free, MONT) in the presence of Fe(II) and reported the formation of monomeric U(IV) as the most plausible sorbed reaction product rather than the commonly observed uraninite (UO<sub>2</sub>). By combining U and Fe speciation (following XANES and Mössbauer analyses) in this study, we suggest an U(VI) reduction mechanism by considering specific site sorption of Fe(II) on MONT. The Fe(II) reactivity related to the surface of Fe-poor montmorillonite (2.6 wt.% Fe-MONT) and nontronite (25.6% NAu-2) was compared to that of

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88 Fe-free montmorillonite (0 wt.% MONT) probing U(VI) as a redox sensitive species under 89 similar experimental conditions. All the experiments were performed in a CO<sub>2</sub>-free atmosphere to compare our results with those obtained in presence of CO<sub>2</sub> by Tsarev et al.<sup>15</sup> and also to 90 91 exclude the effect of carbonate on uranium mobilization. The anoxic (O<sub>2</sub>-free) condition should 92 preserve the oxidation state of ferrous iron and prevent the re-oxidation of reduced uranium 93 species. A high ionic strength (0.05M CaCl<sub>2</sub>) was used to exclude Fe(II) sorption at cation 94 exchange sites, and thus to take into account only for the reactivity of Fe(II) sorbed specifically 95 in the reduction process.

96

#### 97 Materials and Methods

#### 98 Chemicals

99 All the solutions were prepared with boiled, argon (99.9992%) purged Millipore Milli-Q water. 100 NaOH and HCl stock solutions were made from Titrisol ampoules. The Fe(II) stock solutions 101 were freshly prepared from analytical grade FeCl<sub>2</sub>, 4H<sub>2</sub>O (Fluka Chemica), after transferring the 102 required amount to a glove box (JACOMEX) under N<sub>2</sub> atmosphere (CO<sub>2</sub>, O<sub>2</sub> <1 ppmv), 103 dissolved in deoxygenated water (dissolved  $O_2 < 1 \text{ mgL}^{-1}$ ) and acidified with 0.1 M HCl to pH <2 to avoid oxidation.<sup>33 57</sup>Fe(II) stock solutions were prepared by dissolving  $\sim 100$  mg  $^{57}$ Fe(0) in 104 105 concentrated HCl at 100 °C and subsequent dilution with deoxygenated water in glove box. 106 U(VI) solutions were prepared from 1000 mgL<sup>-1</sup> UO<sub>2</sub>(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O (Alfa Aesar) standard 107 solution.

108 Clay minerals

109 The synthetic Fe-free montmorillonite (MONT) (Na<sub>0.30</sub>[(Al<sub>1.70</sub>Mg<sub>0.30</sub>)Si<sub>4</sub>O<sub>10</sub>(OH,F)<sub>2</sub>],nH<sub>2</sub>O) was 110 identical to that used in previous studies.<sup>19,21,25</sup> The synthetic structural Fe(III) bearing 111 montmorillonite (Fe-MONT, 2.6 wt.% Fe) has the representative chemical formula,  $Na_{0.66}(Si_{8.0})$ 112  $(Al_{3.00}Mg_{0.66}Fe_{0.34}III)$ . The natural iron-rich nontronite (NAu-2, 25.8 wt.% Fe)  $[Na_{0.72}(Si_{7.55}Al_{0.50})]$  $Fe_{0.29}^{III}$  (Mg<sub>0.05</sub>Fe<sub>3.54</sub><sup>III</sup>)] was obtained from the Source Clay Minerals Repository, South 113 Australia.<sup>34</sup> Approximately 1% calcite impurities in this clay was removed by using Na-114 115 acetate/acetic acid solution (pH 5) verified by XRD (not shown). All three clays in their Ca-116 exchanged forms were obtained by repeated saturation with 0.05 M aqueous solution of CaCl<sub>2</sub> 117 and clay suspensions were then extensively argon-purged and transferred to the glove box.

#### 118 **Preparation of Fe(II) sorbed smectites**

To evaluate the reactivity of sorbed Fe(II) on smectites, three clay suspensions  $(10.6 \pm 0.2 \text{ gL}^{-1})$ were reacted with  $1.2 \pm 0.2 \text{ mM}$  Fe(II) at pH  $6.0 \pm 0.2$  for 72 h before U(VI) addition. The Mössbauer samples were prepared in a similar way using <sup>57</sup>Fe(II). The experimental conditions are given in Table 1.

#### 123 Kinetic redox experiments

Fe(II)–U(VI) redox kinetic experiments in the presence of three Ca-exchanged smectite clays (MONT, Fe-MONT and NAu-2) were carried out in a closed reactor at room temperature ( $24 \pm 1$ °C) with 0.05 M CaCl<sub>2</sub> ionic background as described in our previous study for MONT.<sup>25</sup> The control samples (without Fe) containing clay and U(VI) was prepared in 50 ml vials by batch method (Table 1).

129

#### 130 <sup>57</sup>Fe Mössbauer spectrometry

<sup>57</sup>Fe Mössbauer experiments were performed at 77K in transmission geometry with a 925 MBq <sup>7</sup>-source of <sup>57</sup>Co/Rh mounted on a conventional constant acceleration drive. The samples containing about 5 mg Fe.cm<sup>-2</sup> were prepared and sealed in a glove box, and then transferred in a bath cryostay. The hyperfine structures were fitted using the MOSFIT program<sup>35</sup> involving quadrupolar components with lorentzian lines; the isomer shift values are referred to that of  $\alpha$ -Fe at RT. The velocity of the source was calibrated using  $\alpha$ -Fe as the standard at RT.

137 X-

#### X-ray absorption spectroscopy

138 X-ray Absorption Near-Edge Structure (XANES) and Extended X-ray Absorption Fine-139 Structure (EXAFS) spectra were collected at the Rossendorf Beamline at the European 140 Synchrotron Radiation Facility (ESRF, Grenoble, France). The energy of the X-ray beam was 141 tuned by a doubled crystal monochromator operating in pseudo channel-cut mode using a 142 Si(111) crystal pair. Two platinum-coated Si mirrors before and after the monochromator were 143 used to collimate the beam into the monochromator and to reject higher harmonics. Uranium L<sub>III</sub>-144 edge spectra were collected in fluorescence mode using a 13-element high purity germanium 145 detector (Canberra) together with a digital signal processing unit (XIA Xmap). Spectra were 146 collected at 15K using a closed cycle He cryostat with a large fluorescence exit window 147 (CryoVac). The energy was calibrated using the maximum of the first derivative of the yttrium 148 K-edge (17038 eV). Between eight and ten scans were recorded for each sample. Dead time 149 correction of the fluorescence signal, energy calibration and averaging of the individual scans were performed with the software package SixPack.<sup>36</sup> Normalization, transformation from 150 151 energy into k space, subtraction of a spline background was performed with WinXAS using 152 routine procedures.<sup>37</sup> Statistical analyses of XANES and EXAFS spectra was performed by Iterative Target-test Factor Analysis (ITFA) using the ITFA software package.<sup>38-40</sup> 153

#### 155 **Results and Discussion**

#### 156 Sorption of Fe(II) and U(VI) on clays

157 The experimental conditions with results from wet chemical analyses are shown in Table 1. 158 Among three clays, NAu-2 has the highest affinity for Fe(II) sorption followed by Fe-MONT 159 and MONT (Figure 1). The sorption of Fe(II) at pH 6 is attributed to cation exchange as well as site-specific sorption to clay edge sites.<sup>21,26</sup> In the absence of Fe(II), the extent of U(VI) sorption 160 161 is less on Fe-MONT than MONT or Nau-2. The kinetic experiments (Fig. 1A & 1B) show a fast 162 uptake of >80% of U(VI) from solution by the three clays with sorbed Fe(II) within 1 hr. The 163 EXAFS results identified surface precipitation of soddyite-like phase only on MONT with 164 sorbed Fe(II) after 72 h, a precipitate which was not observed for Fe-MONT and NAu-2. The 165 removal of aqueous U(VI) by Fe(II) sorbed on MONT is therefore attributed primarily to a fast 166 sorption of U(VI), followed by a slow precipitation of soddvite-like phases probably favored by 167 dissolution of clay which will be discussed in the next section. In Fe(II) sorption experiment on 168 NAu-2, a visible color change is observed from brown to pale green clearly different from 169 commonly observed the deep green color of dithionite reduced NAu-2 (Figure S1 in supporting 170 information). This color change is attributed to inter-valence charge transfer (IVCT) between 171 sorbed Fe(II) and structural Fe(III) reported by other investigations.<sup>15, 28, 29</sup>

#### 172 Speciation of uranium in presence of Fe(II) sorbed on clays

#### 173 XANES and EXAFS

174 The normalized XANES spectrum of the U(VI) reference sample (NAu-2 + U(VI) at pH 6.2 i.e. N1) (Table 1) in Figure 2A shows the characteristic features of the uranyl ( $U^{VI}O_2^{2+}$ ) structure, 175 176 including a white line peak centered at 17179 eV and a shoulder centered at approximately 177 17190 eV. The feature at 17190 eV has previously been shown to result from multiple scattering 178 resonances of the linear uranyl ion structure with the short U-O<sub>ax</sub> bonds.<sup>42</sup> The XANES spectra 179 of M1, M2, M3, F1, F2 and N2 samples are very similar to the U(VI) reference, implying that 180 the dominating phase of uranium in these samples is in +VI oxidation state. This visual 181 observation of the XANES spectra was further statistically verified with ITFA and a U(IV) 182 reference spectrum, showing no coincidence. Therefore, no reduction of U(VI) by Fe(II) sorbed 183 on three clays was observed after 72 h. ITFA analysis of the k<sup>3</sup>-weighted EXAFS spectra 184 revealed the presence of two structurally different components or species. One of the 185 components, prevailing in samples M2 and M3, could be identified as samples an amorphous, 186 soddyite [(UO<sub>2</sub>)<sub>2</sub>SiO<sub>4</sub>.2H<sub>2</sub>O]-like surface precipitate forming in presence of Fe(II) on MONT. 187 Using the spectrum of M1 as representative of the most likely pure U(VI) sorption complex, and 188 that of a soddyite reference as representative of the surface precipitate, we could derive by 189 Iterative Target Test the quantitative speciation of both components in the other sample spectra 190 (Table 2). Only the M series starting after 24 h and increasing with reaction time showed 191 evidence of surface precipitation. For samples F2 and N2, no reduction of U(VI) to lower 192 valence state occurred suggesting that sorbed Fe(II) probably loses its reactivity while being 193 involved in intervalence charge transfer (IVCT) with structural Fe(III) present in tetrahedral 194 and/or octahedral sites for Fe-MONT and NAu-2 clays before reacting with aqueous U(VI).

To focus more on the kinetics of surface precipitation, we analyzed three samples M2a, M2b and
M2c reacted at different time intervals. The XANES and EXAFS quantification (Table 2, 3)

197 shows that neither reduction of U(VI) nor surface precipitation occurred (Table 1) before 24 h in 198 case of M2a and M2b. However, a partial reduction about 21% of total sorbed U(VI) was 199 indicated by XANES in M2c (Table 3). However, the spectral characteristic of M2c with its 200 absence of U-U backscattering is not similar to that of  $U_3O_8(s)$  with mixed U valence used as 201 reference (not shown). Although we cannot unequivocally prove the presence of U-Al or U-Fe 202 backscattering paths most likely masked by the strong multiple scattering peak at 2.9 Å 203 (uncorrected for phase shift), the spectral features of M2c are in line with the formation of an 204 inner-sphere complex like  $\equiv$ Fe(II)-O-U(VI) at the "reduced strong" edge sites coupled to a slow 205 reduction step attributed to the surface oxidation of Fe(II) at the "oxidised strong" edge sites of 206 MONT. No further increase in U(VI) reduction is observed before 72 hr. Moreover, in samples 207 M2 and M3 we did not observe any U(VI) reduction probably due to the fact that surface 208 precipitation starts between 24 h and 72 h and is clearly visible after 72 h (Figure 2 & 3). The 209 total Si concentrations in solution in samples M2a, M2b and M2c show a slight upward trend up 210 to 24 h due to the dissolution of MONT which, however, decreases in sample M2 due to surface 211 precipitation of an uranium silicate-like soddyite solid phases (Table 3). So it is possible that the 212 major part of the signals comes from the U(VI) bearing soddyite-like mineral phase showing no 213 reduction in M2 and M3.

#### 214 Speciation of sorbed Fe on clays with/out U(VI)

#### 215 Mössbauer analyses

The 77K Mössbauer spectra of Fe(II) sorbed on MONT before and Figure 4B after U(VI) addition are compared in Figure 4. The corresponding refined values of the hyperfine parameters are listed in Table 4. According to the values of isomer shift, among four doublets, D1, D2 and 219 D3 are unambiguously assigned to Fe(II) species while D4 to Fe(III) species. The spectrum in 220 Figure 4A must be deconvoluted into four quadrupolar doublets corresponding to (a) 221 exchangeable FeCl<sup>+</sup>, (b) exchangeable Fe(II), (c) Fe(II) sorbed on strong reduced site2 and (d) Fe(II) sorbed on strong oxidized site of MONT.<sup>21</sup> The presence of Fe(III) (D4 doublet) in sample 222 223 M suggests that 10% of total sorbed Fe(II) is oxidized to Fe(III) by "oxidized strong edge sites" 224 of MONT surface before U(VI) addition. This finding is well consistent with that previously found by Charlet et al.<sup>19</sup> and Gehin et al.<sup>21</sup> The spectrum shown in Figure 4B must be 225 226 deconvoluted into at least five doublets. The new doublet D5 with higher quadrupole splitting 227 value is assigned to the Fe(III) sorbed on strong reduced site2 which appeared to the detriment of 228 the oxidation of sorbed Fe(II) of the "strong reduced" site2 of MONT after addition of U(VI). 229 Thus the additional 6% oxidation of Fe(II) involves "strong reduced edge sites" concerned by the 230 U(VI) reduction. On the basis of this result, two mechanisms of U(VI) reduction are proposed.

#### 231 Mechanistic approaches of U(VI) sorption-reduction by Fe(II)/MONT

Our previous study<sup>25</sup> has shown that reduced uranium may exist on MONT and results in surface 232 monomeric U(IV) complex species also shown in recent studies.<sup>15</sup> Therefore, U(VI) reduction 233 234 mechanism is suggested by considering U(VI) and monomeric U(IV) surface complexes formed 235 with Fe(II) specifically sorbed on oxidized and reduced strong sites of MONT. Furthermore, 236 Mössbauer analyses of solid phase Fe speciation (Table 4) reveals that the reduction of U(VI) 237 occurs in two steps: (1) a slow, diffusion controlled process involving H<sub>2</sub>-like species formation resulting from the pre-oxidation of Fe(II) on oxidized strong sites of MONT<sup>19</sup> and (2) a relatively 238 239 faster electron transfer to U(VI) through inner-sphere complexation between U(VI) and Fe(II) 240 sorbed on "reduced" strong sites of MONT.

241 **Mechanism 1**. U(VI) reduction into monomeric U(IV) sorbed species by surface  $H_2$  species 242 produced due to pre-oxidation of Fe(II) specifically sorbed on MONT oxidized strong 243 sites.<sup>19,21,42</sup>

244 
$$Fe^{2+}(aq) + s^{1} \equiv^{3+}(OH^{-})_{3} \rightarrow s^{1} \equiv^{3+}(OH^{-})(O^{2-})_{2}Fe^{2+} + 2H^{+}$$
 (1)

245 
$$s^{1} \equiv^{3+} (OH^{-})(O^{2-})_{2} Fe^{2+} + 2H_{2}O \leftrightarrow s^{1} \equiv^{3+} (O^{2-})_{3} Fe^{3+} (OH^{-})_{2} \dots (H_{2})_{0.5} + 2H^{+}$$
 (2)

246 
$$s^{1} \equiv^{3+} (O^{2-})_{3} Fe^{3+} (OH^{-})_{2} + 2s^{1} \equiv^{3+} (O^{2-})_{3} Fe^{3+} (OH^{-})_{2} \dots (H_{2})_{0.5} + 2U^{VI}O_{2}^{2+} + 2H_{2}O \rightarrow$$

247 
$$(s^{1}\Xi^{3+}(O^{2-})_{3}Fe^{3+}(OH^{-})_{2})U^{VI}O_{2}(OH)_{2} + (s^{1}\Xi^{3+}(O^{2-})_{3}Fe^{3+}(OH^{-})_{2})_{2}U^{IV}(OH)_{2} + 2H^{+}$$
 (3)

248 Mechanism 2. U(VI) reduction by Fe(II) sorbed on reduced strong sites of MONT by inner249 sphere complex formation.

250 
$$Fe^{2+}(aq) + s^2 \equiv^{3+}(OH^{-})_3 \rightarrow s^2 \equiv^{3+}(OH^{-})(O^{2-})_2 Fe^{2+} + 2H^+$$
 (4)

251 
$$2s^2 \equiv {}^{3+}(OH^-)(O^{2-})_2Fe^{2+} + U^{VI}O_2{}^{2+} + 4H_2O \rightarrow (s^2 \equiv {}^{3+}(O^{2-})_3Fe^{3+}(OH^-)_2)_2U^{IV}(OH)_2 + 4H^+$$
 (5)

where  $s^{1}\equiv^{3+}(OH^{-})_{3}$  and  $s^{2}\equiv^{3+}(OH^{-})_{3}$  represent one mole of MONT oxidized and reduced strong sites respectively at pH 6. Equation (3) is attributed to a diffusion controlled process and is therefore slow (days).

#### 255 Precipitation of Soddyite-like mineral phase on Fe(II) sorbed on MONT surface

M2 and M3 suspensions are slightly oversaturated with respect to soddyite (saturation index
0.37) and precipitation may occur (Table 2) according to:

258 
$$2U^{VI}O_2^{2+} + H_4SiO_4 + 2H_2O = (U^{VI}O_2)_2SiO_4 \cdot 2H_2O + 4H^+$$
 (6)

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259 Previously it has been shown that redox potential of mineral/electrolyte suspension is useful in 260 predicting end product reactivity of mineral when reacted with a contaminant.<sup>15,43</sup> In the present 261 study, the redox potential of Fe(II) sorbed on Fe-MONT or NAu-2 suspension is lower than 262 Fe(II)/Fe(III) in solution (Figure S2 in supporting information). Moreover, the amount of Fe(II) 263 sorbed on these clays is nearly 10 times higher than the amount of sorbed U(VI), 5 times higher 264 than stoichiometry would require for complete reduction to occur. However U(VI) reduction is 265 not complete. In MONT 16% of the total sorbed Fe(II) is found to be active in electron transfer 266 to U(VI). On the other hand, Fe(II) sorbed on Fe-MONT and NAu-2 shows stronger affinity to 267 reduce structural Fe(III) before U(VI) addition and therefore loses its reactivity towards U(VI) 268 reduction. One may form "dead-end" – Fe(III)-O-U(VI) complexes at both of these clay surfaces.

269 Working at higher concentration of clay, Fe(II) and U(VI) than those we previously used,<sup>25</sup> 270 we do not show significant impact on the extent of U(VI) reduction but we observe a 271 precipitation of soddyite-like phases on MONT. However, such precipitation is not observed in 272 other systems containing either U(VI)-Fe(III)-bearing clays or Fe(II)-U(VI)-Fe(III)-bearing 273 clays. It can therefore be concluded that the mobility of U(VI) is controlled by a coupled 274 sorption-reduction and uranium silicate mineral phase precipitation reaction in the U(VI)-Fe(II)-275 MONT system whereas only redox-inactive sorption prevails when U(VI) is co-adsorbed on 276 Fe(II)-Fe-MONT and Fe(II)-NAu-2 systems.

277 Environmental implications

The long-term stability of uranium present in nuclear waste disposal sites or bomb shells must be carefully assessed using MONT, Fe-MONT and NAu-2 considering that (1) mobility of U(VI) is mainly controlled by sorption-reduction on Fe(II) sorbed MONT surface coupled with the precipitation of soddyite-like solids at higher U concentrations and (2) the reductive immobilization of U(VI) is not operative in Fe(II)-U(VI)-Fe-MONT and Fe(II)-U(VI)-NAu-2 systems due to inter-valence charge transfer.

284

#### 285 ACKNOWLEDGMENT

This research was funded by ANDRA (project no. #20072484). We are thankful to Dr. Jocelyne Brendlé for synthesis of Fe-MONT and Dr. Laura Leone for Mössbauer analysis of Fe-MONT. Delphine Tisserand and Florian Molton are acknowledged for their technical support within the Analytical Chemistry Platform at ISTerre (OSUG), the ESRF Radioprotection Group and the scientists of the Rossendorf Beamline (ROBL) for providing support during the XAS measurements.

292

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·	Sample		[Fe(II)] <sub>0</sub>	pH <sub>i</sub> _	[Fe(II)] <sub>sorb</sub>	[U(VI)] <sub>0</sub>	рНi	pH <sub>f</sub> _U	[U(VI)] <sub>sorb</sub>
	No.	Clay type	$(\text{mmolL}^{-1})$	Fe	(mmolkg <sup>-1</sup> )	(mmolL <sup>-</sup> 1)	_U		(mmolkg <sup>-</sup> 1)
	Mc		1.24	6.0	20.5	-	-	-	-
	M1	Fe-free (0 wt%)	-	-	-	0.13	6.0	6.3	12.1
	M2 <sup>c</sup>	montmorillonite	1.15	5.9	19.7	0.13	6.0	6.1	12.0
	M3	(MONT)	1.21	6.0	20.6	0.13	6.0	6.0	11.8
	U1 <sup>b</sup>		0.69	6.2	20.0	0.04	6.1	5.5	8.3
	F	Fe-poor (2.6	1.50	5.9	87.3	-	-	-	-
	F1	wt%) montmorillonite	-	-	-	0.11	5.8	5.7	9.5
	F2	(Fe-MONT)	1.50	5.9	85.7	0.13	6.0	5.7	8.8
	N	Fe-rich (25.8	1.08	6.0	103.3	-	-	-	-
	N1	wt%) nontronite	-	-	-	0.13	6.2	5.1	11.8
	N2	N2 (NAu-2)		6.0	103.5	0.14	5.9	5.3	10.4
425 426	<sup>a</sup> [cla	y] =10.6 $\pm$ 0.2 gL <sup>-1</sup> ,	M & U = m	ontmori	illonite (MONT	T); F = Fe-mo	ontmori	llonite	(Fe-MONT)
427	an	d N = Nontronite (N	Au-2); equil	ibration	time of Fe(II)	with clay bef	ore U(V	VI) add	ition = 72 h
428	an	d equilibration time	of U(VI) afte	er Fe(II)	sorption on cla	y = 72 h.			
429	<sup>b</sup> stuc	ly by Chakraborty et a	al. <sup>25</sup>						
430	<sup>c</sup> sam	ples prepared using <sup>5</sup>	<sup>7</sup> Fe(II)						

424	TABLE 1.	Experimental	conditions	of sorption	of U(VI	) on Fe(II	) sorbed ont	o smectite clays <sup>a</sup>
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433	TABLE 2.	Quantification	of U(VI)	precipitation	molar fraction	as soddyite-like	mineral phase
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434 and U(VI) sorbed as obtained by ITFA of XANES (12.10–12.25 keV).

			U(VI) precipitated	U(VI) sorbed	
Sample No	рН	t (hr)	as soddyite-like phase	onto Fe(II)	Sum
			(UO <sub>2</sub> ) <sub>2</sub> SiO <sub>4</sub> .2H <sub>2</sub> O	sorbed on clays	
Soddyite	-	-	1.00	0.00	1.00
M1	6.0	72	0.00	1.00	1.00
M2a	5.8	0.16	0.02	0.97	0.99
M2b	5.7	1	0.05	0.92	0.98
M2c	5.5	24	0.08	0.87	0.95
M2	6.0	72	0.26	0.51	0.77
M3	6.0	72	0.75	0.00	0.75
F1	5.8	72	0.08	0.89	0.97
F2	6.0	72	0.03	0.92	0.95
N1	-	-	0.00	1.00	1.00
N2	5.9	72	0.04	0.99	1.03

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439 TABLE 3. Quantification of U oxidation states (molar fraction) by ITFA of XANES (12.10-

No         pH         t (III) $0(VI)$ $0(IV)$ Sum $\mu molL^{-1}$ $\mu molL^{-1}$ M1         6.0         72         1.00         0.00         1.00         -         -           M2a         5.8         0.16         0.94         0.06         0.99         345         11           M2b         5.7         1         0.95         0.05         1.00         351         20           M2c         5.5         24         0.79         0.21         1.00         358         18           M2         6.0         72         0.91         0.09         1.00         335         3           U1a         6.1         72         0.87         0.14         1.01         -         -	Sample	aIJ	t (lam)			Sum	[Si] <sub>solution</sub> in	[U] <sub>solution</sub> in
M1 $6.0$ $72$ $1.00$ $0.00$ $1.00$ $ -$ M2a $5.8$ $0.16$ $0.94$ $0.06$ $0.99$ $345$ $11$ M2b $5.7$ $1$ $0.95$ $0.05$ $1.00$ $351$ $20$ M2c $5.5$ $24$ $0.79$ $0.21$ $1.00$ $358$ $18$ M2 $6.0$ $72$ $0.91$ $0.09$ $1.00$ $335$ $3$ U1 <sup>a</sup> $6.1$ $72$ $0.87$ $0.14$ $1.01$ $-$	No	рн	t (nr)	U(VI)	$U(\mathbf{IV})$	Sum	µmolL <sup>-1</sup>	µmolL <sup>-1</sup>
M2a       5.8       0.16       0.94       0.06       0.99       345       11         M2b       5.7       1       0.95       0.05       1.00       351       20         M2c       5.5       24       0.79       0.21       1.00       358       18         M2       6.0       72       0.91       0.09       1.00       335       3         U1 <sup>a</sup> 6.1       72       0.87       0.14       1.01       -       -	M1	6.0	72	1.00	0.00	1.00	-	-
M2b       5.7       1       0.95       0.05       1.00       351       20         M2c       5.5       24       0.79       0.21       1.00       358       18         M2       6.0       72       0.91       0.09       1.00       335       3         U1 <sup>a</sup> 6.1       72       0.87       0.14       1.01       -       -	M2a	5.8	0.16	0.94	0.06	0.99	345	11
M2c       5.5       24       0.79       0.21       1.00       358       18         M2       6.0       72       0.91       0.09       1.00       335       3         U1a       6.1       72       0.87       0.14       1.01       -       -	M2b	5.7	1	0.95	0.05	1.00	351	20
M2       6.0       72       0.91       0.09       1.00       335       3         U1a       6.1       72       0.87       0.14       1.01       -       -	M2c	5.5	24	0.79	0.21	1.00	358	18
U1 <sup>a</sup> 6.1 72 0.87 0.14 1.01	M2	6.0	72	0.91	0.09	1.00	335	3
<sup>a</sup> Study, by Chakraborty at al <sup>25</sup>	U1 <sup>a</sup>	6.1	72	0.87	0.14	1.01	-	-
- Study by Chaklabolity et al	M2 U1ª <sup>a</sup> Study by C	6.0 6.1 Chakraborty	72 72 v et al. <sup>25</sup>	0.91	0.09	1.00	-	-
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440 12.25 keV) and solution data (pH, [Si], [U] in  $\mu$ molL<sup>-1</sup>).

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451 TABLE 4. Refined values of <sup>57</sup>Mössbauer hyperfine parameters estimated from the spectra
452 presented in Figure 4.

Sample No.	Site		$\delta \pm 0.02$	Γ± 0.02	$\begin{array}{c} \Delta E_Q \pm \\ 0.02 \end{array}$	RA ± 2
	Fe(II)_exch_FeCl <sup>+</sup>	D1	1.37	0.41	3.40	36
Μ	Fe(II)_exch_Fe <sup>2+</sup>	D2	1.32	0.34	3.03	33
(Fe(II)/MONT)	Fe(II)_strong_red2	D3	1.28	0.34	2.62	21
	Fe(III)_strong_ox	D4	0.47	0.49	0.58	10
	Fe(II)_exch_FeCl <sup>+</sup>	D1	1.37	0.28	3.42	39
	Fe(II)_exch_Fe <sup>2+</sup>	D2	1.34	0.38	3.04	32
M2C	Fe(II)_strong_red2	D3	1.30	0.32	2.59	13
(re(II)/MONT+O(VI))	Fe(III)_strong_ox	D4	0.47	0.46	0.58	10
	Fe(III)_strong_red2	D5	0.48	0.46	1.24	6
	Fe(III)_strong_red2	D4 D5	0.47	0.40	1.24	6

 $\delta$  (mm s<sup>-1</sup>) isomer shift with respect to metallic  $\alpha$ -Fe(0) at 300K;  $\Delta E_Q$  (mm s<sup>-1</sup>) quadrupole

455 splitting; RA (%) relative abundance,  $\Gamma$  (mm s<sup>-1</sup>) full-width at half-height.

#### 462 FIGURE CAPTIONS

- 463 Figure 1. Kinetics of U(VI) sorption on (A) Fe(II)/MONT; (B) Fe(II)/Fe-MONT, Fe(II)/NAu-2.
- 464 [U(VI) added to Fe(II)/clay suspensions at t = 0].
- 465 Figure 2. U L<sub>III</sub>-edge XAS spectra of U(VI) sorbed on MONT, Fe-MONT and NAu-2 in the
- 466 presence of Fe(II) compared to the U(VI) and soddyite references, (A) fitted XANES; (B)
- 467 EXAFS and (C) Fourier transform of EXAFS.
- 468 Figure 3. U L<sub>III</sub>-edge XAS spectra of kinetics of U(VI) sorption on MONT in the presence of
- 469 Fe(II) compared to the U(VI) (M1 sample in Table 1) and U(IV) colloid references, (A) fitted
- 470 XANES; (B) EXAFS and (C) Fourier transform of EXAFS.
- 471 Figure 4. <sup>77</sup>K Mössbauer spectra of <sup>57</sup>Fe(II) sorbed on MONT (A) before; (B) after U(VI)
  472 addition.
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Figure 2



Figure 3



Figure 4