Uranium Redox Transformations after U(VI) Coprecipitation with Magnetite Nanoparticles

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ABSTRACT: Uranium redox states and speciation in magnetite nanoparticles coprecipitated with U(VI) for uranium loadings varying from 1000 to 10 000 ppm are investigated by X ray absorption spectroscopy (XAS). It is demonstrated that the U M₄ high energy resolution X ray absorption near edge structure (HR XANES) method is capable to clearly characterize U(IV), U(V), and U(VI) existing simultaneously in the same sample. The contributions of the three different uranium redox states are quantified with the iterative transformation factor analysis (ITFA) method. U L₃ XAS and transmission electron microscopy (TEM) reveal that initially sorbed U(VI) species recrystallize to nonstoichiometric UO_{2+x} nanoparticles within 147 days when stored under anoxic conditions. These U(IV) species oxidize again when exposed to air. U M₄ HR XANES data demonstrate strong contribution of U(V) at day 10 and that U(V) remains stable over 142 days under ambient conditions as shown for magnetite nanoparticles containing 1000 ppm U. U L₃ XAS indicates that this U(V) species is protected from oxidation likely incorporated into octahedral magnetite sites. XAS results are supported by density functional theory (DFT) calculations. Further characterization of the samples include powder X ray diffraction (pXRD), scanning electron microscopy (SEM) and Fe 2p X ray photoelectron spectroscopy (XPS).

INTRODUCTION

Uranium is the main constituent of spent nuclear fuel (SNF), but it can be also found in high quantities in actinide (An) contaminated sites.¹ Developing the safety case for the safe disposal of radioactive waste requires mechanistic under standing of the interaction of the waste products with repository components. In case of water accessing the waste container, radionuclides will react with the corrosion products, which potentially represent a very relevant reactive barrier retaining pollutants release in the repository near field. Of particular interest is the detailed understanding of An interactions with iron (Fe) oxides, for example, magnetite (Fe₃O₄) considered as a corrosion product of Fe based container materials.² A number of laboratory studies have been performed to clarify the fate of uranium in such systems. Due to the complex redox processes induced by Fe(II)/Fe(III) species, uranium is often found to exist in a mixture of redox states. Uranium has two main environmentally relevant redox states, U(IV) and U(VI). U(V) is believed to form as an intermediate redox species and exhibits a poorly understood geochemical behavior. The only evidence of U(V) in nature is the uranium mineral wyartite.³ Whereas U(VI) and U(IV) are usually found, depending on conditions, as a result of microbial and Fe(III)/ Fe(II) driven redox processes, $^{4-10}$ only a few studies report U(V) as a relevant redox species. In earlier studies U(V) was detected after Fe(II) catalyzed transformation of U(VI) ferrihydrite to goethite (α FeO(OH))¹¹ and in different Fe (oxyhydr)oxides phases where Fe₃O₄ was considered as one of the possible phases to stabilize U(V).¹² Also Nico et al. reported on the possibility of U(V) or U(VI) incorporation into the octahedral position of the α FeO(OH)/Fe₃O₄ after ferrihydrite remineralization.¹³ Later studies attempted to specifically detect U(V) in different Fe systems under controlled conditions with the aim to reconsider the relevance of U(V) for uranium containing (geo)chemical systems.^{14–17} In most of the reported investigations, X ray absorption spectros copy (XAS) based methods, that is, U L_3 X ray absorption near edge structure (XANES) and extended X ray absorption fine structure (EXAFS) as well as U 4f X ray photoelectron spectroscopy (XPS) are used to investigate the U speciation and redox states. XANES spectra are sensitive to the U coordination environment and redox states but the spectra are dominated by broad features due to large core hole lifetime broadening effects.¹⁸ This challenges the characterization and quantification of the different U redox states, especially when U(V) is also present in the material. On the other hand, the XPS technique can successfully quantify mixed U redox states if there is sufficiently high uranium content (>1000 ppm) in the sample.^{12,14,16,19} However, a clear identification of U(V) species can be hampered by low signal to noise ratio of the spectra for low uranium contents. XPS has high surface sensitivity and is performed in ultrahigh vacuum, which can lead to potential changes of the samples. To handle these challenges, complementary highly sensitive techniques are needed for thorough characterization of the U redox states in such systems. The high energy resolution XANES (HR XANES) method at the U M_{4.5} absorption edges was demonstrated to be very valuable for studies related to U redox state analysis in complex uranium systems containing mixed U redox states.^{20,21} The U M₄ HR XANES experiments need to be performed in He environment comprising the sample, analyzing crystals and detector in order to avoid loss of intensity by scattering and absorption of photons in air. Unlike XPS, vacuum conditions are not necessary. The samples can be investigated in the form of solids, wet pastes, suspensions or liquids. U M_{4.5} HR XANES takes a great advantage over the conventional U L₃ XANES. Reduced core hole lifetime broadening results in better resolved spectral features allowing more precise redox state analysis. Due to the dipole selection rule $(\Delta l = \pm 1)$ the electrons are excited with high probability from U $3d_{3/2,5/2}$ to U 5f unoccupied states. Therefore, the U M₄ HR XANES technique is a direct probe of the unoccupied U 5f valence states, which play a significant role in the chemical bonding of the An elements. 20-24

In the present study we investigated the U redox states and speciation in the final product formed by coprecipitation of U(VI) with Fe₃O₄ for variable U loadings (1000–10 000 ppm). The U L₃/M₄ HR XANES and U L₃ XANES/EXAFS as well as transmission electron microscopy (TEM) are applied. Scanning electron microscopy (SEM), X ray powder diffraction (pXRD), Fe 2p_{3/2} XPS, geochemical calculations and density functional theory (DFT) are used to characterize the U containing Fe₃O₄ samples. The main aim is to verify the presence of U(V) in this system. A long term study of samples kept under anoxic (up to 480 days) and oxidizing conditions (up to 226 days) was performed to elucidate the stability of the potential U(V) species in the Fe₃O₄ nanoparticles. We demonstrate that the applied U M₄ HR XANES method is capable of detecting U(IV), U(V), and U(VI) at relatively low (1000 ppm) total U concentrations being present simultaneously in the same sample. The contributions of the three different U redox states are quantified with the iterative transformation factor analysis (ITFA) method.²⁵

MATERIALS AND METHODS

Preparation of the Samples. Milli Q H₂O was used for the preparation of all samples. To remove dissolved O₂ and CO₂ Milli Q H₂O was bubbled for several hours with Ar outside and then for at least 1 h inside an argon (Ar) glovebox. Uranium containing Fe_3O_4 nanoparticles were synthesized by direct precipitation^{26–28} inside an Ar glovebox equipped with pH, $E_{\rm h}$ electrodes and a dropping funnel using the following procedure: to a 50 mL Teflon container the calculated amounts of aqueous FeCl₃, FeCl₂ and an aliquot of aqueous $U(VI)O_2Cl_2$ (1000, 3000, 6000, and 10 000 ppm U: Um1 Um10 samples) were added followed by dropwise addition of 0.5 M NaOH (Backer, CO₂ free) to pH 7.5-8.0. The suspension was stirred overnight, pH and E_h values were recorded and pH was adjusted if necessary by adding 0.1 M NaOH. All samples were then stored in the form of suspension in the Ar glovebox. The UO_2 (several μm particle size), U_4O_9 (used from^{20,29}) and 3000 ppm U(VI) adsorbed onto maghemite (γ Fe₂O₃, spectra are named Umh) serve as reference compounds. The γ Fe₂O₃ was prepared by heating freeze dried magnetite nanoparticles at 200 °C during 2 h in air.³⁰ 3000 ppm U as U(VI)O₂Cl₂ was then adsorbed on the γ Fe_2O_3. More than 99% of U(VI) was adsorbed after 50 days as determined with inductively coupled plasma mass spectrometry (ICP MS). Experimental details for U, Fe and salt concentrations as well as pH, $E_{\rm h}$ measurements are given in Supporting Information (SI) Table S1.

For the U L_3/M_4 HR XANES experiments, solids were separated from supernatant using a Nd magnet. The super natant was decanted and the solids were used in the form of a wet paste. All samples were prepared in an Ar glovebox at 1-2ppm of O₂ level and less than 1 ppm of CO₂. An inert gas Plexiglas sample holder comprising a double containment (two separated 10 μ m polypropylene films) has been designed and used for the spectroscopic measurements. To avoid contact of the samples with air they were transported 1 day prior to the experiments in a gastight aluminum cylinder filled with Ar and opened very shortly prior to the experiments.

For the U L_3 XAS experiments, subsamples Um1 Um10 from the same batch were separated using a Nd magnet after 147 days of aging in the Ar glovebox. After these experiments the Um1 sample was placed in a nonhermetically closed plastic vial and kept under ambient, aerobic conditions (Um1a sample). The Um1a sample was investigated with the U M_4 HR XANES (142 days in air, total 289 days) and with the U L_3 XAS (XANES and EXAFS) techniques (226 days in air, total 373 days).

X-ray Absorption Spectroscopy (XAS). U L_3/M_4 HR XANES experiments were performed at the ID26 beamline, ESRF, Grenoble, France³¹ (Um1 Um10) and at the INE Beamline, ANKA, Karlsruhe, Germany²² (Um1a) with Johann type X ray emission spectrometers.^{24,32,33} The U L_3 XAS (XANES and EXAFS) experiments were performed in fluorescence mode using a five element Ge solid state

fluorescence detector (Canberra) at the INE Beamline. The ATHENA and ARTEMIS program parts of the IFFEFIT program package were used for data reduction and analyses of the EXAFS spectra.^{34,35} Details on the experiments, the quantitative U oxidation states analyses of the U M_4 HR XANES spectra with the ITFA method for the 10 day aged samples and the EXAFS analyses are given in the Supporting Information.

Additional Characterization Methods. High resolution TEM (HR TEM) and SEM images were recorded with a FEI Tecnai G2 F20 X TWIN instrument operated at 200 kV and a FEI Quanta 650 FEG ESEM, respectively. pXRD patterns were measured with a laboratory based Bruker AXS D8 powder diffractometer and at the SUL X Beamline (ANKA, Karlsruhe, Germany) (SI).³⁶ The FIT2D and DIFFRAC.EVA V3.1 programs were used to analyze the data.³⁷

DFT Calculations. The experiments are supported by DFT calculations utilizing plane wave basis sets with periodic boundary conditions as implemented in the Vienna Ab initio Simulation Package (VASP).³⁸⁻⁴⁴ Further details are given in SI.

Details on the thermodynamic calculations and the XPS experiments are reported in SI. Table S2 comprises the description of the studied samples and the applied character ization techniques.

RESULTS AND DISCUSSION

Characterization of the Fe_3O_4 Nanoparticles. The formation of the Fe_3O_4 nanoparticles can be described with reaction 1,

$$\mathrm{Fe}^{2+} + 2\mathrm{Fe}^{3+} + 8\mathrm{OH}^{-} \to \mathrm{Fe}_{3}\mathrm{O}_{4} \downarrow + 4\mathrm{H}_{2}\mathrm{O} \tag{1}$$

Measured E_h pH values for samples containing 1000 ppm U (Um1) and 10 000 ppm U (Um10) are located inside the stability field of Fe₃O₄ according to the Fe Pourbaix diagram (SI Figure S1A); redox conditions of the samples comprising 3000 ppm U (Um3) and 6000 ppm U (Um6) are placed at the border of $Fe_3O_4 - FeO(OH)$ (cr) (goethite) stability fields.⁴⁵ $E_{\rm h}$ pH conditions are such ($E_{\rm h}$ = -210 to -390 mV; pH 7.5-8.0) that subsequent reduction of U(VI) to U(IV) would be expected according to the U Pourbaix diagrams (SI Figures S1B, C). SEM images (SI Figure S2) and powder XRD (pXRD) patterns (SI Figure S3) reveal that crystalline Fe₃O₄ nanoparticles with octahedral shape and a size of 15-40 nm are formed. Energy dispersive X ray analysis (EDX) does not reveal segregated uranium precipitates. No goethite or any other Fe containing crystalline phases are found. The Fe(II) content is slightly below that of stoichiometric Fe_3O_4 (Fe(II)/Fe_{TOT} = 0.33) at Fe(II)/Fe_{TOT} = 0.28 \pm 3% for all samples (Um1 Um10) aged for 310 days as indicated by the reduced intensity of the Fe(II) $2p_{3/2}$ peak of the Fe 2p XPS spectra (SI Figure S4). There is no clear trend in the level of Fe(II) oxidation as a function of the U(VI) concentration. In view of the low U/Fe mass ratios (0.0014-0.014), this is indeed not to be expected.

Redox States of U in the Fe₃O₄ Nanoparticles Aged for 10 days. We characterized the U redox state in the Um1 Um10 samples aged in inert atmosphere for 10 days with the U L₃ and M₄ HR XANES spectroscopy techniques. The U L₃ and M₄ HR XANES spectra of the Um1 Um10 samples are depicted in Figure 1A and B, respectively.

The X ray absorption spectra typically shift to higher energies by increasing the U redox state due to the reduced



Figure 1. From bottom to top: U L₃ HR XANES (A) and U M₄ HR XANES (B) spectra of the Umh reference (U(VI) sorbed on maghemite for 55 days) and the Um10, Um6, Um3, Um1 samples aged for 10 days in anoxic conditions as well as the U_4O_9 and UO_2 references.

screening of the $2p_{3/2}$ (L₃)/3d_{3/2} (M₄) core hole by the decreased electronic charge density on the U atoms. Smaller energy shifts of about ± 0.5 eV can be induced by variations of the electronegativity of the bonding partner, changes in symmetry, short and long range atomic order, etc., for the same U redox state as suggested by U L₃ XANES studies.^{46,47} One exception of the general trend are the U L₃ XANES spectra of U(V) and U(VI) forming short (<1.9 Å) axial bonds with two O atoms $(UO_2^+/UO_2^{2+}, U(V)/U(VI) \text{ yl})$; the maxima of the most intense absorption resonance (white line, WL) of the U(V) yl and UO_2 spectra have similar energy positions. The WL of the U(VI) yl spectrum is only slightly shifted to higher energy and it can potentially have the same energy position as e.g. U(V) species with more symmetric coordina tion environments associated with loss of short uranyl bonds, that is, uranate type of U(V) species. This effect is induced by the large electronic density in the vicinity of U due to the strong covalent bond of U(V)/U(VI) with the two axial O atoms. Such anomalous behavior is not observed for U M_{4.5} HR XANES spectra. Energy shifts within a range of ± 0.1 eV can be caused by differences in short and long range atomic order around the absorbing atom as for example observed for different UO_3 phases, but the spectra shift to higher energies for higher U oxidation states.^{21,48} For example the energy shift between UQ₂ and U(V) yl ($[U(V)O_2(CO_3)_3]^{5-}$) is ~1.2 eV, whereas between U(V) yl and U(VI) yl ($[U(V)O_2(CO_3)_3]^{5-}$) is ~1.0 eV.⁴⁹

The WL of the U L₃ HR XANES spectrum of the Um10 sample is located at similar energy position as the WL of the spectrum of Umh. U has oxidation state U(VI) in the Umh sample. In addition, the feature marked with line c clearly visible in the U L₃ HR XANES Umh and Um10 spectra (Figure 1A) is characteristic for the U(VI)O₂⁻²⁺/U(V)O₂⁺ entity. These results suggest predominant U(VI) yl contribution in the Um10 sample. The absorption resonance marked with line c becomes less intense in the spectra of the Um1 sample. These spectra suggest that the Um1 sample if at all contains low amount of uranyl species. In addition, the WL and the post edge absorption resonance marked with line c' become broader

and asymmetric from the Um10 to the Um1 spectrum. Such asymmetric WLs have been previously described as being characteristic for uranate type of U(V)/U(VI) species.^{19,46,50,51} However, it can not be excluded that this U species dominant in the Um1 sample has redox state (IV). The local symmetry of these U(V)/U(IV) species should be rather different compared to the U(V)/U(IV) in U₄O₉ since there is no resemblance between the Um1 and the U₄O₉ spectra. It is clear that the U L₃ HR XANES spectra are more sensitive to small structural and oxidation states changes compared to the conventional U L₃ XANES.^{18,52} But the method is still limited by large core hole lifetime broadening effects and does not allow to reliably identify and quantify the U(IV), U(V), and U(VI) anticipated to be simultaneously present in the samples.

Additional information on the U redox states is obtained by applying the U M₄ HR XANES method. Figure 1B depicts the U M₄ HR XANES spectra of the Um1 Um10 samples and UO₂, U₄O₉ as well as Umh for comparison. U₄O₉ contains U(IV) and U(V) visible by the two main peaks separated by ~1 eV.²⁰ These spectral features are named d and e in the U M₄ HR XANES spectrum of U₄O₉ (Figure 1B). The main absorption peak f of the Umh spectrum is shifted by ~0.4 eV to higher energies as compared to peak e and it is assigned to U(VI). Spectral features f' and f'' are also typical for U(VI) yl ions with short (~1.77–1.91 Å) trans dioxo bonds.^{23,48,53} These features exist for U(V) yl species but are considerably shifted (f' ~ 0.6 and f'' ~ 2.5 eV) toward the main peak.⁴⁹

Much more clear compared to the U L_3 HR XANES, the U M_4 HR XANES spectrum obtained for the Um1 sample demonstrates major contributions of U(V) (peak e) and minor fraction of U(IV) (peak d) (Figure 1B). It is also evident that the intensity of feature d increases while moving from the Um10 to the Um1 spectrum, the energy position of feature e shifts to lower energies and features f' and f'' lose intensity. These spectral changes imply that the relative contribution of U(VI) decreases, whereas the U(IV) content rises going from 10 000 ppm U (Um10) to 1000 ppm U (Um1) in the samples (Figure 1B).

Quantification of the U Redox States. We performed quantitative analysis of the U M4 HR XANES spectra by the ITFA method.²⁵ ITFA is used to decompose the spectral mixtures into the spectra and the fractions of the components (SI Figure S5). The experimental spectra can be sufficiently reproduced (SI Figure S5A) by using linear combinations of the three ITFA extracted component spectra (SI Figure S5B), so that the residual is similar to the expected experimental error. Only the first three eigenvectors show a signal while the eigenvectors 4, 5, and 6 do not contribute to the data (SI Figure S5C), hence only three components are necessary to describe the variations in the spectral mixtures. The extracted component spectra corespond to the U(IV), U(V), and U(VI) redox states. The analysis shows that the Um1 and Um10 samples contain 19% U(IV), 81% U(V) and 1% U(IV), 62% U(V), 37% U(VI), respectively (SI Table S3). The U M₄ HR XANES technique and the ITFA approach are clearly capable of detecting and quantifying the three different redox states: U(IV), U(V), and U(VI), being present simultaneously in the same sample.

Redox States and Local Atomic Environment of U in the Fe_3O_4 Nanoparticles Aged for 147 days. We performed U L₃ XAS (XANES and EXAFS) studies for the magnetite suspensions containing U stored for 147 days in anoxic conditions. For technical reasons, the redox states of U in the Um1, Um3, Um6, and Um10 samples could not be analyzed with the U M_4 HR XANES method 147 days after the preparation of the samples.

The U L_3 XANES spectra of the Um10 Um1 samples are depicted in Figure 2A. The energy positions of the WLs and the



Figure 2. From bottom to top: U L_3 XANES (A) and FT EXAFS spectra in R space (B) of the UO₂ reference and the Um10, Um6, Um3, Um1 samples aged for 147 days in anoxic conditions as well as the Umh reference (U(VI) sorbed on maghemite for 330 days). The FT EXAFS and the best fit to the data are given in colored rhombs and in dash dot, respectively.

general shapes of the spectra of the Um10 sample and UO₂ are very similar. We conclude that the initial U(VI) yl has reduced mainly to U(IV) in the Um10 sample. The spectra of the Um10 Um1 samples exhibit a trend and are shifted up to ~1.5 eV to higher energies compared to UO₂ going from the Um10 to the Um1 sample (lines a and b in Figure 2A). The WL becomes broader and the postedge absorption resonance at ~17 218 eV transforms from a single asymmetric (c') to a broad peak shifted to higher energies observed also for the U L₃ HR XANES spectra of the 10 days aged samples (Figure 1A). We obtain additional insights from the comparison of the conventional U L3 XANES spectra measured for the Um1 samples aged for 10 and 147 days (SI Figure S6). The U L₃ XANES for the Um1 sample aged for 147 days is shifted ~2.5 eV to lower energies compared to the spectrum of the sample aged for 10 days, which is an evidence for higher U(IV) content (Figure S6). This result implies that reduction of U(V) to U(IV) has continued for the Um1 sample 10 days after its preparation. We assume that the higher oxidation state of U in the Um1 sample aged for 147 days is the U(V) species found for the 10 days aged Um1 sample. U(VI) was not found for the 10 days aged Um1 sample by the ITFA U M₄ edge HR XANES analyses and since the Um1 Um10 samples were handled in the same way, i.e. anoxic and reducing conditions, it is likely that U has reduced in all samples. Therefore, the observed energy shift can not be explained with major U(IV) and minor U(VI)contributions.

Alternatively, if the U has completely reduced to U(IV), this can be U(IV) in a different coordination environment compared to the U(IV) typical for the Um10 sample. However, published spectra for U(IV) in different coordination environ ments show no significant differences in the energy positions of the WL maxima of the U L_3 XANES spectra.⁵⁴

The FT EXAFS spectra and their best fits for the Um1, Um3, Um6 and Um10 samples after 147 days aging are depicted in Figure 2B (see also SI Figures S7–10). The first coordination sphere of U is best modeled with three U–O distances (R). R(U-O1) = 1.69(2) 1.73(2) Å is an untypically short bond length for U(VI) O_{axial}, which has been previously observed and controversially discussed.^{55,56} Particularly Conradson et al. proposed that these short U–O distances are characteristic for U(VI) in a series of UO_{2+x} compounds.⁵⁵ The structural parameters obtained from the fit to the FT EXAFS are similar with and without modeling these peaks; we chose not to consider these unusually short U–O distances (<1.75 Å) in our structural model since they can be also part of a background signal. More detailed discussion on their potential structural meaning is not in the scope of our study.

Due to the low signal to noise ratio of the EXAFS data particularly for the Um1 and Um1a samples we have performed and report fits to the spectra for k range until 9.5 Å⁻¹ (SI Figure S10). The main U–O2 distance obtained from the fits to the data continuously decrease from 2.26(1) Å to 2.18(2) Å within the Um10 Um1 series, whereas the U-O3 distance remains within the range R = 2.41(1) Å (Table S4, Figure S11). The coordination numbers (N) vary: N(U-O1) = 0.2(1) 0.4(2), N(U-O2) = 2.2(2) 3.0(3) and N(U-O3) = 3.2(3) 1.8(2) (SI Table S4, Figure S11). The observation that the first O coordination sphere of U splits into two dominant peaks was reported for UO_{2+x} (x = 0-0.2) compounds where the split became more pronounced for large x.⁵⁵ We therefore might assume that U in our samples exists mainly as non stoichiometric UO_{2+x} containing mixed U redox states. However, the trend for these two main U-O distances is reversed compared to our EXAFS results since R O2 and R O3 grow as a function of the increasing nonstoichiometric oxygen in UO_{2+x} with R(U-O2) = 2.22-2.26 Å and R(U-O3) =2.36–2.42 Å. 55 This can be explained by the presence of a second U site in our samples in addition to U(IV)/U(V)/U(VI) constituents of UO_{2+x} particles. This U species becomes more relevant at low uranium concentration (1000 ppm U), which agrees with the results from the U L₃ XANES data. The R(U-O2) = 2.18 Å for the Um1 sample is within the range R =2.05–2.20 Å reported in the literature as typical for U(V)compounds.¹² These EXAFS results do not contradict to the assumption that the U(V) found in the Um1 10 days aged sample is at least partially preserved. This potential U(V)species is likely coordinated by Fe as the best fits to the FT EXAFS spectra reveal. This is based on the finding that coordination of U to Fe atoms is more prominent for the Um1 compared to the Um10 sample. Two U-Fe interatomic distances are resolved: N(U-Fe1) = 0.6(3), R(U-Fe1) =3.13(3) Å in Um10 and N(U-Fe1) = 4.5 ± 1.7 , R(U-Fe1) = 3.19(1) Å in Um1. A U–Fe2 shell with a minor contribution (N = 0.5(3) 1.0(8)) is modeled too (SI Table S4). Somewhat longer R(U-Fe) (+ 0.04 Å) and N(U-Fe) = 6 (fixed) values have been reported for U incorporated into Fe octahedral sites.15

We still need to consider possible U(IV) species incorporated in Fe₃O₄. Kerisit et al. calculated similar R(U– Fe1) = 3.18 Å for U(V) and R(U–Fe1) = 3.19 Å for U(IV) incorporated in Fe₃O₄. The R(U–O) = 2.10 Å is much shorter for U(V) compared to R(U–O) = 2.23 Å for U(IV) and closer to the experimental results for the oxidized Um1a sample with R(U–O) = 2.13 Å (cf. section Redox States of U in the Fe₃O₄ Nanoparticles Exposed to Air).⁵⁷ As we discuss below the U(V) species in Um1 and Um1a are very similar but there is less interference with near U–O shells for the Um1a compared to the Um1 samples since UO_2/UO_{2+x} is oxidized to form U(VI) yl in the former. To the best of our knowledge there is no report for U(IV) incorporated in octahedral Fe₃O₄ sites. Based on these evidences, we conclude that U(V) not U(IV) is more likely to be incorporated in the structure of Fe₃O₄. (cf. DFT calculations).

In contrast, UO_{2+x} species dominate in the Um10 sample. The R(U-O2) = 2.26 Å and R(U-O3) = 2.41 Å obtained from the EXAFS analyses for this sample are very similar to the reported values for $UO_{2.17} UO_{2.20}$.⁵⁵ The authors describe strong variations of the R(U–O) in the first coordination sphere as a function of x in UO_{2+x} . In our system R(U-O2)changes as a function of the U concentration due to the increasing contribution of the second U(V) species, whereas R(U-O3) remains constant. Hence it is apparent that the proposed UO_{2+x} has similar stoichiometry in all samples. Presence of UO_{2+x} in the Um1 Um10 samples is also demonstrated by the intense peak at about 3.84 Å characteristic for the scattering of the photoelectron from U atoms in the second U coordination sphere. The N(U-U1) vary within 5.8(8) 2.4(8) in the Um10 Um1 series. The coordination numbers are significantly smaller than those found for bulk UO_2 : N(U-O) = 8 and N(U-U) = 12. This can be due to structural disorder and the large contribution of U surface atoms in the small nanoparticles, resulting in destructive interference and thus in decreasing amplitudes of the EXAFS.^{41,48} However, it can be also explained by the fraction of incorporated U atoms versus UO_{2+x} nanoparticles. There is a distinct trend in the coordination numbers. The N(U–U1) and N(U-O3) decrease, whereas N(U-Fe1) increases going from Um10 to Um1 (SI Figure S11). We assume that minor redox changes have occurred for the Um1 sample since the characteristic for UO_{2+x} : N(U-O3) = 1.8(2), N(U-U1) =2.4(8) and for U incorporated in magnetite: N(U-Fe1) = 4.5 \pm 1.7 correspond to about 19% of U(IV) and 81% U(V) found also from the quantitative analyses of the U M4 HR XANES data for 10 days aging of the samples. Correspondingly, if the about 37% U(VI) found for the Um10 sample aged for 10 days has reduced to U(IV) and if we take into account the initial 1% U(IV), the coordination numbers will be similar to those reported for the 147 day aged Um10 sample: N(U-O3) = 3.2, N(U-U1) = 5.8. The coordination numbers obtained from EXAFS analyses can have uncertainties up to 30%. But since we compare them in a system with systematic structural variations, the magnitude of the uncertainty is significantly reduced.

TEM analyses clearly confirm our interpretation of the EXAFS data since it detects crystalline UO₂ nanoparticles with size of about 4–5 nm (Figure 3A) possibly grown on the surface of the Fe₃O₄ particles (Figure 3B) with lattice parameters very close to stoichiometric UO₂. Due to the relatively large uncertainty for the obtained lattice parameter ($a = 0.271 \pm 0.004$ nm) it is not possible to distinguish between UO₂ and UO_{2+x}. Considering also the EXAFS results, we conclude that they are both likely.

It is useful to discuss potential formation of U(IV), U(V) or U(VI) inner sphere complexes sorbed on Fe₃O₄. Indeed U(IV) inner sphere complexes stable within a few months have been reported to form on Fe₃O₄ surfaces; the R(U–O) is elongated and the N(U–O) is higher (R(U–O) = 2.42 Å, N(U–O) = 9) compared to UO₂ (R(U–O) = 2.35 Å, N(U–O) = 8).⁵⁴ For this U(IV) sorbed species almost no U–U coordination is



Figure 3. HR TEM image of UO_2 nanocluster (blue box) and a Fe_3O_4 nanoparticle (magenta box) in the Um1 (1000 ppm U) sample aged for 330 days in anoxic conditions (A). pXRD pattern of the Um1 sample aged for 480 days in anoxic conditions (top) and a reference pXRD pattern of Fe_3O_4 (ICSD 26410) (bottom) (B). U(V) incorporated into an octahedral iron site into the Fe_3O_4 structure as optimized by DFT; O atoms are in red, Fe atoms in brown, U atoms in blue.

found similarly to U(IV) sorbed onto a Ti substituted Fe₃O₄. Moreover R(U–Fe) = 3.56–3.59 Å are reported, which are significantly longer compared to the distances found in our system, R(U–Fe1) = 3.13–3.19 Å. Indeed the R(U–O) = 2.42 Å is comparable to the R(U–O3) for our Um10 Um3 samples (SI Table S4). However, our EXAFS analyses report higher N(U–U) for the Um10 sample, N(U–U1) = 5.8, which is in a good agreement with the crystalline UO₂/UO_{2+x} particles found by TEM. Latta et al. showed that UO₂ nanoparticles, are formed for high U surface coverage for samples aged for several months under anoxic conditions.⁵⁴ They found systematic change of the only one U–O and the U–U distances as a function of the U concentration not observed for our system. We conclude that formation of crystalline UO₂/UO_{2+x} nanoparticles is much more probable compared to sorbed U(IV) species.

The possibility for U(V) stabilized on Fe(II) containing mineral surfaces as an inner sphere complex has been discussed;⁵⁸ however no exact structural model was described. The stabilization of U(V) seems to be more likely in the form of mixed U(V)–U(VI) oxyhydroxide.⁵⁹ For this compound two very long R(U–O_{axial}) = 2.41–2.44 Å and four shorter equatorial O ligands R(U–O_{equat}) = 2.06 Å are found. Our EXAFS results do not report U–O bond distances, which correspond to the distance obtained for the equatorial ligands, therefore this model does not seem to be appropriate.

U(VI) complexes sorbed on Fe oxides have two axial O atoms at about 1.79 Å, six equatorial O atoms at about 2.38 Å and N(U–Fe) \sim 2(5), R(U–Fe) > 3 Å.⁶⁰ Since we find N(U–

Fe) \sim 4.5 and no strong evidence for stabilization of uranyl, we conclude that the formation of U(VI) yl is not likely.

DFT Calculations. The aim of the calculations is to verify if it is energetically favorable to incorporate U(V) into a magnetite site. In the DFT calculations with periodic boundary conditions we used a $2 \times 2 \times 2$ super cell of magnetite and replaced one Fe^{2+} and one Fe^{3+} by U^{5+} and one vacancy $(Fe^{2+}+Fe^{3+} \rightarrow U^{5+}+\Box)$. The Fe^{2+} and Fe^{3+} were chosen to be close to each other, hence the presence of the vacancy allows the substitution of Fe by U to relax more easily. Since U(V) is an open shell system we carefully monitored the occupation of the 5f orbitals and the orientation of the spin of this electron. U(V) can be incorporated into a Fe²⁺ or Fe³⁺ octahedral site, since both cases are energetically very close to each other (cf. Figure 3C). We found that the orientation of the spin is important. The spin of the 5f electron points in the opposite direction compared to the spins of the two replaced Fe ions. For the R(U-O2) and the R(U-Fe1) we calculated theoretical values of 2.15(3) Å and 3.18(4) Å, respectively, which are in excellent agreement with the experimental results, R(U-O2) =2.18 Å and R(U-Fe1) = 3.18 Å, respectively (SI Table S4).

Redox States of U in the Fe₃O₄ Nanoparticles Exposed to Air. In order to verify possible existence of U(V) in octahedral Fe₃O₄ environment the Um1 sample aged for 147 days in Ar atmosphere was exposed for 142 days (U M₄ HR XANES) and for additional 84 days (U L₃ XANES and EXAFS) to air. The U M₄ HR XANES spectrum of Um1a (142 days in air) is depicted in Figure 4A. The shoulder characteristic for



Figure 4. U M_4 HR XANES spectra of the Um1 and Um1a samples as well as the Umh reference. The Um1 sample was aged for 10 days. The Um1a sample is the Um1 sample aged for 147 days exposed to air for additional 142 days (289 days total aging time) (A). FT EXAFS of the Um1 and Um1a samples. The Um1a sample is the Um1 sample aged for 147 days exposed to air for additional 226 days (373 days total aging time) (B).

U(IV) is not visible, whereas features f' and f" typical for U(VI) yl appear in the spectrum. The main peak maximum has an energy position very similar to the spectrum of the Um1 sample aged for 10 days in Ar, which confirms that the main redox state of U is U(V) (Figure 4A). After exposing this Um1a sample to additional 84 days in air U L₃ XAS (XANES and EXAFS) spectra were recorded. The U L₃ XANES is not as informative as the U M₄ HR XANES but it can help to verify if any substantial changes of the U oxidation state has taken place.

The energy positions of the WL and the post edge absorption resonance of the U L_3 XANES spectra of the Um1 aged for 10 days and the Um1a (226 days in air) samples are comparable, which is an indication that the majority of the U(V) remains stable in the Um1a sample exposed to air for additional 84 days (SI Figure S6).

The FT EXAFS spectra of the Um1a (226 days in air) and the Um1 (aged for 147 days in Ar) samples are compared in Figure 4B. It is apparent that the peaks characteristic for U coordination to O and Fe are preserved: N(U-O2) = 2.8(2), R(U-O2) = 2.13(1) Å; N(U-O3) = 0.8(2), R(U-O3) =2.36(2) Å and N(U–Fe1) = 6.2 ± 1.7 , R(U–Fe1) = 3.19(1) Å, whereas the peak describing the coordination to U is absent in the FT EXAFS Um1a spectrum. The U(IV) in the UO_{2+x} particles is likely oxidized to $U(VI)O_2^{2+}$ as suggested from the R(U-O1) = 1.78(1) Å with N(U-O1) = 0.7(1). The disappearance of the peak characteristic for U at about 3.84 Å can be explained by increased disorder during the partial oxidation of U leading to destructive interference of the scattered photoelectron waves. Alternatively, the U-U coordination can disappear due to a phase transformation to a U(VI) phase, which does not contain U atoms at this distance. We did not find crystalline U containing phases for the Um1a sample by applying pXRD. The relatively long term stability (142 days) of U(V) upon exposure to air (Um1a) can be explained by its incorporation into the octahedral sites of Fe₃O₄, where it is possibly protected against oxidation.

U Redox Transformation Model. We propose the conceptual model presented in the abstract graphics. More than 90% of initially added U(VI) has adsorbed, most likely on ferrihydrite^{16,61} which is formed in the first stage after Fe(III) titration by NaOH solution and later recrystallizes to Fe₃O₄ through intermediate Fe (oxyhydr)oxide phases.²⁷

Part of U(VI) stays adsorbed onto the Fe₃O₄ after 10 days of aging time. We suppose that the U(V) species have incorporated into the structure of Fe₃O₄ during the formation of the nanoparticles as suggested by the more symmetric local coordination environment compared to uranyl and coordination to Fe. Minor formation of U(IV) in the form of UO₂/UO_{2+x} or sorbed onto the surface is also possible.

After 147 days the U(VI) species have undergone phase transformation and reduction to UO_2/UO_{2+x} , whereas major part of the incorporated U(V) species is preserved. When exposed to air for more than 142 days U(IV) oxidizes to U(VI), whereas U(V) remains stabilized incorporated into octahedral sites of Fe₃O₄.

The experimental results illustrate that reduction of U(VI) to U(IV) is likely to take place, expected from geochemical calculations, with apparently the highest U(IV) fraction for the sample with the highest uranium content (10 000 ppm). The relative fractions of U(IV), U(V) and U(VI) depend on the initial uranium concentration while the kinetics of the uranium redox processes depends on the pH/redox conditions, Fe(II)/ Fe_{TOT} ratio as well as the rate of electron transfer between Fe²⁺ and Fe³⁺ ions in the octahedral sublattice.

The present study demonstrates the capability of U M₄ HR XANES to clearly distinguish three different U redox states, U(IV), U(V), and U(VI), in one sample and shows also the potential to identify redox states for other actinides. Previously, U(V) was observed in different Fe(III) (oxyhydr)oxides.^{11–17} Our observation that U(V) can also be stabilized in Fe₃O₄, which is a mixed Fe(II)/Fe(III) phase, suggests that the U(V) stabilization range can be extended to more reducing

conditions. When exposed to air, apparently, the surrounding mineral matrix is able to protect U(V) against oxidation. This might explain the observation of significant U(V) fractions in redox systems where predominantly U(IV) or U(VI) would be expected considering thermodynamic calculations. This shows that there is a lack of kinetical data to reliably describe and predict uranium speciation in such complex systems. Further studies of U interaction with Fe phases with varying Fe(II)/Fe(III) ratios will be of interest extending the redox/pH conditions where U(V) can be stabilized.

Notably for the discussion on the relevance of U(V) species in the environment, U $M_{4,5}$ HR XANES combined with EXAFS and other techniques appears to be a powerful spectroscopic tool, which is able to provide distinct answers. Another aspect of future studies is to analyze systems with lower, environ mentally relevant U concentrations, for example, <100 ppm U, and to investigate a representative U–Fe containing natural system with a varying set of geochemical conditions leading to different U redox and immobilization/mobilization scenarios.

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