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Influence of Uranyl Speciation and Iron Oxides on Uranium Biogeochemical Redox Reactions

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Abstract

Uranium is a pollutant of concern to both human and ecosystem health. Uranium’s redox state often dictates its partitioning between the aqueous- and solid-phases, and thus controls its dissolved concentration and, coupled with groundwater flow, its migration within the environment. In anaerobic environments, the more oxidized and mobile form of uranium (UO$_2^{2+}$ and associated species) may be reduced, directly or indirectly, by microorganisms to U(IV) with subsequent precipitation of UO$_2$. However, various factors within soils and sediments may limit biological reduction of U(VI), inclusive of alterations in U(VI) speciation and competitive electron acceptors. Here we elucidate the impact of U(VI) speciation on the extent and rate of reduction with specific emphasis on speciation changes induced by dissolved Ca, and we examine the impact of Fe(III) (hydr)oxides (ferrihydrite, goethite and hematite) varying in free energies of formation on U reduction. The amount of uranium removed from solution during 100 h of incubation with *S. putrefaciens* was 77% with no Ca or ferrihydrite present but only 24% (with ferrihydrite) and 14% (no ferrihydrite) were removed for systems with 0.8 mM Ca. Imparting an important criterion on uranium reduction, goethite and hematite decrease the dissolved concentration of calcium through adsorption and thus tend to diminish the effect of calcium on uranium reduction. Dissimilatory reduction of Fe(III) and U(VI) can proceed through different enzyme pathways, even within a single organism, thus providing a potential second means by which Fe(III) bearing minerals may impact U(VI) reduction. We quantify rate coefficients for simultaneous dissimilatory reduction of Fe(III) and U(VI) in systems varying in Ca concentration (0 to 0.8 mM), and using a mathematical construct implemented with the reactive transport code MIN3P, we reveal the predominant influence of uranyl speciation,
specifically the formation of uranyl-calcium-carbonato complexes, and ferrihydrite on the rate and extent of uranium reduction in complex geochemical systems.
Introduction

A century of uranium-ore mining and processing have created an inventory of uranium and co-contaminants in the environment that threaten human and ecosystem health. Uranium’s partitioning between the solid and aqueous phases, and thus potential for transport, in surface and subsurface environments is controlled in part by its oxidation state. In oxidized, carbonate-bearing settings, uranium commonly exists as the uranyl ion (UO$_2^{2+}$) associated with a variety of uranyl-carbonato complexes such as UO$_2$(CO$_3$)$_2^{2-}$ and UO$_2$(CO$_3$)$_3^{4+}$ (Lovley, 1991; Clark et al., 1995; Brooks et al., 2003;), which, in the absence of Ca, account for ~95% of U(VI) in solution in a typical groundwater (Abdelouas et al., 1998a) —a pH of ~6.9, temperature of 16°C, and HCO$_3^-$ and U(VI) concentrations of 1 mM and 4 μM respectively. Inclusion of Ca, however, results in two ternary calcium-uranyl-carbonato species, CaUO$_2$(CO$_3$)$_3^{2-}$ and Ca$_2$UO$_2$(CO$_3$)$_3$, emerging as dominant aqueous species, often accounting for >99% of U(VI) in solution (Bernhard et al., 1996, 2001; Kalmykov and Choppin, 2000; Kelly et al., 2003; Dong and Brooks, 2006).

In anaerobic environments, U(VI) is often reduced to U(IV) in the presence of dissimilatory metal-reducing bacteria (DMRB) (Lovley, 1991; Gorby and Lovley, 1992; Liu et al., 2002) and sulfate reducing bacteria (SRB) (Ganesh et al., 1997; Payne et al., 2002) that couple the reduction of metals such as U(VI) with the oxidation of carbon or H$_2$; in the specific case of U(VI) reduction, the result subsequent to reduction is precipitation of sparingly soluble UO$_2$ (Lovley, 1991; Brooks et al., 1999; Fredrickson et al., 2000). In fact, microbially mediated reductive immobilization of uranium has been studied as a potential form of stimulated natural attenuation in several contaminated sites (Anderson et al., 2003; Michalsen et al., 2006; Wu et
al., 2006b). However, several factors including competing electron acceptors in the form of nitrate and Fe(III) (hydr)oxides (Abdelouas et al., 1998b; Wielinga et al., 2000; Wu et al., 2006a), for example, and the potential for re-oxidation of UO$_2$ by a variety of oxidants including molecular oxygen and Fe(III) (Wan et al., 2005; Moon et al., 2007; Wu et al., 2007), may limit reductive immobilization and thus diminish the long term effectiveness of this remediation strategy. Additionally, several studies have focused on describing kinetic behavior and quantifying rates of uranium reduction by DMRB and SRB; both Spear et al. (2000) and Lui et al. (2002) reproduced U(VI) reduction trends with first order models, and Luo et al. described concurrent U(VI) and sulfate reduction with a saturation kinetic model (2007). However, changes in uranyl speciation, particularly those driven by the presence of Ca, can drastically impact microbial reduction of uranium (Brooks et al., 2003; Neiss et al., 2007; Stewart et al., 2007). Brooks et al. (2003), for example, observed a 40% decrease in U(VI) reduction by *Shewanella putrefaciens* over a 30 h period upon the addition of 0.45 mM Ca, as compared to reduction without Ca. Additionally, dissolved Ca and calcium-bearing minerals can decrease U(VI) adsorption to soils and sediments, increasing the propensity for migration in the subsurface (Fox et al., 2006; Meleshyn et al., 2009; Stewart et al., 2010).

The presence of Fe(III) imparts a variety of complicating constraints on uranium reduction, and consequently on its mobility under transport conditions. Firstly, uranyl (and associated species) can adsorb to the surface of Fe(hydr)oxide mineral phases (predominately in the absence of Ca); reductive dissolution of Fe(hydr)oxides thus can subsequently release U(VI) into solution, increasing aqueous concentration. Additionally, owing to comparable (and overlapping) redox couples of U (VI/IV) and Fe (III/II), iron can act as an oxidant or reductant of uranium
depending on specific geochemical conditions that often dictate uranyl speciation (Figure 1, 
(Ginder-Vogel et al., 2006)). Similarly Fe(III) can act as a competing terminal electron acceptor 
in microbial respiration, potentially decreasing both the rate and extent of uranium reduction. 
Wielinga et al. (2000) for example, observed a 52% decrease in uranyl reduction by Shewanella 
alga, strain BrY, in the presence of ferricydrite, compared to only uranyl in solution, over a 
period of 10 h, while, in contrast, neither goethite nor hematite impacted uranyl reduction. 
Dissimilatory U(VI) and Fe(III) reduction by DMRB may occur simultaneously due to 
overlapping redox couples and different forms of substrate (aqueous vs. solid), as well as distinct 
enzyme pathways. As summarized previously (DiChristina et al., 2005), Shewanella respiration 
on solid phase Fe(III), including Fe (hydr)oxides, likely proceeds via terminal reductases 
associated with the outer membrane of the cell; the terminal reductase(s) for U(VI) reduction 
appear to reside in the periplasm (for Gram-negative bacteria) and possibly on the outer-
membrane.

The intricate association between iron and uranium combined with the controlling influence of 
uranyl speciation convolutes our ability to quantitatively predict the fate and transport of U in 
soils and sediments. Limiting our ability to predict U reduction is the absence of rate 
information specifically describing the impacts of uranyl speciation, particularly those involving 
ternary complexes of Ca, or soil/sediment matrix effects. The aim of this study is thus to provide 
a quantitative framework on bacterial reduction of U(VI) at varying concentrations of dissolved 
calcium and to determine the impact of Fe(III) (hydr)oxides, varying systematically in free 
energy of formation, on reduction rates. In addition to a kinetic analysis, we develop a 
quantitative biogeochemical model for U transformation using the reactive transport code
MIN3P (Mayer et al., 2002), which is broadly applicable to geochemical problems involving kinetically controlled redox and mineral dissolution/precipitation reactions along with equilibrium hydrolysis, aqueous complexation, ion exchange, and surface complexation reactions. The results of this study provide an understanding of geochemical limitations on microbial uranium reduction imposed by Ca and modifications induced by Fe (hydr)oxides within the mineral matrix of soils and sediments, helping to predict the potential for migration in environmental settings.

**Materials and Methods**

**Cell Culturing and Harvesting**

*Shewanella putrefaciens*, strain CN32, a facultative, dissimilatory metal-reducing bacterium, was used to evaluate the effect of calcium on biological uranyl reduction in batch studies involving ferrihydrite-coated quartz sand. The culture was obtained from the American Type Culture Collection (ATCC catalogue number BAA-453), cultured aerobically in tryptic soy broth (TSB) until late log phase, and then frozen in 20% glycerol at -80°C where it was stored until needed.

*Shewanella putrefaciens* was grown aerobically by placing 1 mL of frozen stock culture in 100 mL of TSB. The culture was placed on a shaker at room temperature for 12 h (late log phase) before it was transferred into the same medium and incubated while shaking for an additional 12 h. It was then centrifuged at 4500 rpm for 12 min, re-suspended in 100 mL of bicarbonate buffer, and centrifuged a second time. Finally, the culture was suspended in 100 mL of PIPES buffered artificial groundwater medium for 30 min until the start of the experiment. The artificial groundwater medium consisted of the following ingredients (in mg/L): KHCO₃, 380;
KCl, 5; MgSO₄, 50; NaCl, 30; NH₄Cl 0.95; KH₂PO₄, 0.95; and 1 mL Wolfe’s mineral solution. The media was buffered with 10 mM PIPES (1,4-piperazinediethanesulfonic acid) at pH 7.0 and made anoxic by boiling and cooling under a stream of O₂-free N₂ gas. Further details are described in Stewart et al. (2007).

**Synthesis of Iron (hydr)oxide-coated Sands**

Ferrihydrite was prepared according to the method described by Brooks et al. (1996). A solution of ferric chloride was rapidly titrated with sodium hydroxide over a period of approximately 10 min until a pH of 7 was reached. Chloride and Na were then removed from the ferrihydrite with sequential rinses.

Goethite was prepared using a slightly modified version of Atkinson et al. (1967). Ferric nitrate was combined with concentrated sodium hydroxide in the absence of CO₂. An effort was made to keep CO₂ out of the product during initial synthesis but not during the dialysis purification stage; CO₂ adsorbed during product synthesis can significantly impact the zero point of charge of the product (Vangeen et al., 1994). Sodium hydroxide was slowly pumped into the ferric nitrate solution over a period of several hours until a pH of 12 was reached during continuous stirring. The slurry was then placed in a 60°C oven for 24 h, and finally salts (and, in particular, residual nitrate) were removed by dialysis for a period of approximately 10 d.

Finally, hematite was prepared following the method described by Schwertmann and Cornell (2000) in which a concentrated ferric nitrate solution is added gradually over a period of 4 h to boiling, distilled water. The solution was stirred constantly. The product was then cooled
overnight and purified by dialysis for approximately 7 d to remove excess salts (in particular nitrate). X-ray diffraction analysis was performed on all three Fe (hydr)oxides to ensure purity of product.

The iron (hydr)oxides were prepared individually and then used to coat quartz sand as reported previously (Brooks et al., 1996; Hansel et al., 2003). Briefly, the iron (hydr)oxide slurry was poured over the quartz sand and mixed thoroughly by hand. The mixture was then left to dry for 48 h before being washed with de-ionized water. Iron concentration on the sands was approximately 10 g/Kg (1% by weight).

**Uranium(VI) Reduction Reactions**

In order to investigate the effects of aqueous calcium concentration on uranium reduction, we assembled batch systems containing *S. putrefaciens*, Fe(III) (hydr)oxide coated sand, artificial groundwater medium, uranyl acetate, and varying concentrations of CaCl₂, under anoxic conditions in a glovebag (Coy Laboratory Products) with a N₂ (95%):H₂ (5%) atmosphere. Lactate was provided in the groundwater medium as a carbon source and electron donor at a concentration of 3 mM. All solutions were made anoxic by boiling and cooling under a stream of N₂ gas. Each 125 mL serum vial contained 1.0 g of Fe (hydr)oxide-coated sand, ≈10⁷ cells of *S. putrefaciens*, and 105 mL of groundwater media. Anoxic stock solutions containing U-acetate, CaCl₂, and KHCO₃ were allowed to equilibrate overnight before being steriley injected into the batch bottles. Each batch system began with 0.168 mM U, added in the form of uranyl acetate, UO₂(C₂H₃O₂)₂, and a Ca²⁺ concentration of either 0 mM, 0.4 mM, 0.6 mM, or 0.8 mM (added as CaCl₂). The batch systems were assembled in a glovebag and then shaken at room
temperature outside the glovebag between samplings; bottles were brought into the glovebag
during sampling. A separate experiment was run for each of the three Fe (hydr)oxides (goethite,
ferrihydrite, and hematite) as well as an experiment with no iron (no-Fe control). A replicate set
of ferrihydrite experiments were conducted for 100 h in order to measure solution and solid
associated Fe(II) for input to the model. All of the aforementioned biotic systems were
conducted in triplicate and error values were calculated as standard deviation of the three
measurements divided by the square root of the number of measurements. Abiotic controls for
each of the iron (hydr)oxides, and one without iron, were conducted in duplicate. The pH of all
experimental systems was 7.0 for the duration of the study.

Aqueous Analyses
Aqueous samples were withdrawn using sterile syringes and filtered through 0.2 μm membranes
inside a glovebag at hourly intervals for the first 8 h of the experiment. Subsequent samples
were collected approximately every 12 to 24 h for the next 7 d (with the exception of the
ferrihydrite systems used for model input that were sampled for 4 d). A final sample was
collected on day 22. All samples were analyzed by inductively coupled plasma-optical emission
spectrophotometry (TJA/IRIS Advantage) to determine the concentration of total uranium, iron,
and calcium in solution. The ferrozine method (Stookey, 1970) was used to measure Fe(II). In
systems containing ferrihydrite that were used for model input it was necessary to determine the
total amount of Fe(II) produced. Ferrozine was also used to extract Fe(II) adsorbed to the surface
of the solid as described previously (Ginder-Vogel et al., 2006). Following aqueous sampling,
groundwater media from each system was decanted and ferrihydrite-coated sand from each
bottle was transferred to a small vial. Five-mL of ferrozine reagent were then added to each 1 g
of ferrihydrite-coated sand and allowed to equilibrate for 20 s, and 2 mL of ferrozine solution were then removed for analysis, allowing for quantification of Fe(II) both in solution and adsorbed to the surface of ferrihydrite.

**Model Development**

**Conceptual Model**

The primary reaction of interest in the system described here is the reduction of aqueous U(VI)$_{(aq)}$ to UO$_2$(s), given by:

$$UO_2^{2+} + H_{2(aq)} \rightarrow UO_2(s) + 2H^+, \quad (1)$$

coupled to the oxidation of lactate,

$$C_3H_5O_3^- + 2H_2O \rightarrow C_2H_3O_2^- + HCO_3^- + H^+ + 2H_{2(aq)}^+, \quad (2)$$

In these simulations, we use the intermediate species H$_2$(aq) to represent electron equivalents as a means of computationally, but not mechanistically, decoupling the oxidation and reduction reactions.

In carbonate-bearing solutions the speciation of the uranyl ion is dependent on the concentration of Ca$^{2+}$ in solution, with the following equilibrium reactions pertaining:

$$(UO_2)_2 CO_3(OH)_3^- + 4H^+ \rightarrow 2UO_2^{2+} + HCO_3^- + 3H_2O \quad K = 10^{-11.19} \quad (3)$$

$$UO_2(CO_3)_3^{4-} + 3H^+ \rightarrow UO_2^{2+} + 3HCO_3^- \quad K = 10^{-9.1470} \quad (4)$$

$$UO_2(CO_3)_2^{2-} + 2H^+ \rightarrow UO_2^{2+} + 2HCO_3^- \quad K = 10^{-4.048} \quad (5)$$

$$UO_2CO_3_{3(aq)}^+ + H^+ \rightarrow UO_2^{2+} + HCO_3^- \quad K = 10^{-0.389} \quad (6)$$

$$Ca_2UO_2(CO_3)_3_{3(aq)}^+ + 3H^+ \rightarrow UO_2^{2+} + 2Ca^{2+} + 3HCO_3^- \quad K = 10^{-0.432} \quad (7)$$
\[ \text{CaUO}_2\left(\text{CO}_3\right)_3^{2-} + 3\text{H}^+ \rightarrow \text{UO}_2^{2+} + \text{Ca}^{2+} + 3\text{HCO}_3^- \quad K = 10^{-5.587} \quad (8) \]

where uranyl-carbonato species are the dominant species in the absence of Ca; in its presence, calcium-uranyl-carbonato species become the dominant species. Equilibrium constants for reactions 3-6 are from Guillaumont et al. (2003) and constants for reactions 7 and 8 are provided by Dong and Brooks (2006). For completeness, several other uranyl-carbonato and uranyl-hydroxide species are included in the simulation; however, these species have only minor affects on the aqueous speciation of the system. In these simulations we have assumed equilibrium conditions governed by the above set of equations.

The rate of U(VI) reduction (Equation 9) is dependent on the speciation of the uranyl ion, where the rate of reduction in solutions dominated by uranyl-carbonato species is much faster than in solutions dominated by calcium-uranyl-carbonato species (Brooks et al., 2003; Stewart et al., 2007). This dependence is described by the rate expression;

\[ R_1 = k_1 \sum \{\text{U}^{(VI)^*}\} \frac{k_1^S}{K_{H_{2(aq)}}^S} \left[ \frac{H_{2(aq)}}{H_{2(aq)}} + \frac{H_{2(aq)}}{H_{2(aq)}} \right] + k_{ib} \sum \{\text{U}^{(VI)^+}\} \left[ \frac{H_{2(aq)}}{K_{H_{2(aq)}}^S} \left( \frac{H_{2(aq)}}{H_{2(aq)}} + \frac{H_{2(aq)}}{H_{2(aq)}} \right) \right] \quad (9) \]

where \( k_1 \) and \( k_{ib} \) are rate constants, the curly brackets represent species activity, \( x_1 \) and \( x_{ib} \) are the order of the rate with respect to \{U(VI)^*\} and \{U(VI)^+\}, respectively, square brackets represent total component concentrations, \( K^S \) is a half saturation constant, and terms U(VI)^* and U(VI)^+ identify the dominant U(VI) species in the absence and presence of calcium, respectively; specifically,

\[ \sum \{\text{U}^{(VI)^*}\} = \left\{\text{UO}_2\left(\text{CO}_3\right)_3(\text{OH})_3\right\} + \left\{\text{UO}_2\text{CO}_3_{(aq)}\right\} + \left\{\text{UO}_2(\text{CO}_3)_2^{2-}\right\} + \left\{\text{UO}_2(\text{CO}_3)_3^{3-}\right\} \quad (10) \]
\[ \sum \{U(VI)^{+}\} = \{Ca_2UO_2(CO_3)_3(aq)\} + \{CaUO_2(CO_3)_3^{2-}\} \]  

(11)

and in practice \( k_1 \) is set much higher than \( k_{1b} \). \( k_1 \) and \( k_{1b} \) are independent of pH and carbonate concentration as these effects are taken into account in the overall U(VI) reduction rate equation (Equation 9) through changes in concentration (and thus activity) of various U(VI) species (Equations 10 and 11).

The rate of lactate oxidation is given by;

\[
R_2 = k_2 \left( \frac{[C_3H_5O_3^-]}{K_{C_3H_5O_3^-} + [C_3H_5O_3^-]} \right) \left( \frac{K_{H_2(aq)}^{I}}{K_{H_2(aq)}^{I} + [H_2(aq)]} \right)
\]

(12)

where \( k_2 \) is the rate constant and specifies the maximum oxidation rate, and \( K^{I} \) is an inhibition constant. The first term provides a rate-dependence on lactate, but in these experiments lactate is provided in large excess. The second term couples the rate of lactate oxidation to the uranium reduction through the component \( H_2(aq) \), which decreases the lactate oxidation rate with increasing \( H_2(aq) \) concentration. This simplified rate expression for lactate oxidation does not take into account microbial growth and decay parameters and assumes that microbial populations are established at the outset of the experiments and do not change significantly through the course of the experiment.

We observed that the inclusion of ferrihydrite to U(VI)/lactate systems decreased the reduction rate of U(VI) in some cases (Stewart et al., 2007) and hypothesized that Fe(III) was competing for lactate. The reductive dissolution of ferrihydrite is given by;

\[
Fe(OH)_3(s) + 0.5H_2(aq) + 2H^+ \rightarrow Fe^{2+} + 3H_2O
\]

(13)

resulting in the production of Fe(II). The reductive dissolution of ferrihydrite is modeled
as a kinetically controlled irreversible reaction:

\[
R_{13} = k_{13} \left( \frac{H_{2(aq)}}{K^S_{H_{2(aq)}} + H_{2(aq)}} \right)
\]  

where the rate is primarily governed by the availability of the electron donor (expressed here as \(H_{2(aq)}\)), provided solid phase ferrihydrite is still present. Note that the rate expressions for U(VI) and Fe(III) reduction (equations 9 and 14) both contain equivalent electron donor terms, and thus are competitively linked to lactate oxidation.

An important aspect of the conceptual representation of the redox system described above is the competition between U(VI) and Fe(III) reduction. This is represented using an electron donor, symbolized here as \(H_{2(aq)}\), that represents any common donor (lactate, acetate, etc…) used in U(VI) and Fe(III) respiration. For the present case using lactate, the total rate of lactate oxidation is controlled by the rate constant \(k_2\), effectively setting the rate of \(H_{2(aq)}\) production, and the competition between the two reduction reactions involving U(VI) and Fe(III) is based on the relative magnitude of the respective rate constants, \(k_1\), \(k_{1b}\), and \(k_{13}\), with the total rate of these reactions dependent on the availability of \(H_{2(aq)}\) (electron equivalents) supplied by the lactate oxidation reaction.

In the simulations, adsorption of U(VI) to ferrihydrite surfaces is accounted for by using the initial aqueous concentrations after equilibration with ferrihydrite. Adsorption of Fe(II) is not considered in the model as experimental results are presented as the sum of both aqueous and adsorbed concentrations. Finally, Ca adsorption was minimal and thus not necessary for inclusion in the simulations.
The conceptual framework presented in this model has some common features with both the Liu et al. (2002) and Luo et al. (2007) models. The first-order Liu et al. (2002) model also assumes a constant microbial population and describes U(VI) reduction with a first-order rate law. The Luo et al. (2007) model incorporates saturation and inhibition terms (arising from nitrate reduction); however, their model accounts for microbial growth and decay which this model does not. Neither of these models, however, captures the effect of U(VI) speciation on reduction rates.

**Model Parameters**

In total, two sets of simulations were run to represent the four no-ferrihydrite experiments and the four with-ferrihydrite experiments. Each set consisted of varying Ca$^{2+}$ concentrations at 0.0, 0.4, 0.6 and 0.8 mM in the form of CaCl$_2$. The simulations included 9 components: H$^+$, UO$_2^{2+}$, HCO$_3^-$, Ca$^{2+}$, lactate, acetate, H$_2$(aq), Fe$^{2+}$, and Fe$^{3+}$; 32 aqueous species were included in the model to accurately determine mineral solubilities. Equilibrium constants for speciation reactions are from the WATEQ4F (Ball and Nordstrom, 1991) and MINTEQA2 (Allison et al., 1991) databases, and are updated where possible to be consistent with the Visual MINTEQ database. Initial solution chemistry was set to represent that of the batch experiments as is described above. In order to simulate the buffering capacity of the batch experiments, the simulations included an artificial buffering reaction to maintain the pH at a value of 7.0.

The simulations were run with a no-flow, arbitrary volume of water with effectively 100% porosity to represent the batch experiments. The simulations with ferrihydrite included a volume
fraction of $4.14 \times 10^{-5} \text{ m}^3 \text{ ferrihydrite/m}^3$ to represent the ratio of ferrihydrite to solution in each batch experiment.

The rate expressions presented above leave 7 adjustable parameters for the non-ferrihydrite simulations and 8 for the with-ferrihydrite simulations, including 4 half-saturation/inhibition constants and 4 rate constants. The lactate half-saturation constant is unimportant, as lactate is assumed to be in excess in both the experiments and simulations. The H$_2$(aq) half-saturation (Equations 9 and 14) and inhibition constants (Equation 12) are used to couple the oxidation and reduction reactions and are both set at $1 \times 10^{-8} \text{ mol L}^{-1}$. This relatively low value ensures that the rates of the reactions remain closely coupled; however, the model results are not sensitive to this value.

Of more importance to the simulation results are the rate constants, particularly, $k_1$, $k_{1b}$, $k_2$, and $k_{13}$. In order to constrain the four key rate constants, these parameters were not adjusted individually for each of the eight simulations, but were kept constant for all simulations. Briefly, $k_1$ was calibrated to measured aqueous uranyl concentrations in the no-calcium/no-ferrihydrite simulations; subsequently, $k_{1b}$ was calibrated in the 0.8 mM calcium/no-ferrihydrite simulations; $k_{13}$ was calibrated to reproduce measured total ferrous iron concentration (adsorbed + aqueous) for the with-ferrihydrite simulations. The maximum rate of lactate oxidation, $k_2$, was determined for the with-ferrihydrite simulations as experimental results have shown that the total amount of lactate oxidation in U(VI) plus Fe(III) systems exceeds that of U(VI)-only systems (Revill, personal commun.). All rate parameters were determined by visual fit to the experimental data.
Results

Uranium(VI) Speciation

Formation of the ternary \( \text{Ca}_2\text{UO}_2(\text{CO}_3)_3 \) complex progressively increases, becoming the dominant uranyl species, over a Ca concentration range of 0 to 1 mM. At a calcium concentration of 0.2 mM, 43% of uranium in solution is associated with one of the uranyl-carbonato complexes at pH 7 with \([\text{CO}_3]_T = 3.8 \text{ mM and } [\text{U(VI)}] = 0.168 \text{ mM} \) (Figure 2)—uranyl-calcium-carbonato complexes account for the remaining 57% of the U(VI) species under these conditions. An increase in Ca concentration to 0.8 mM results in 95% of the uranyl residing within the uranyl-calcium-carbonato complexes. The chemical speciation modeling program Visual MINTEQ was used to calculate these values.

This shift in speciation from a system dominated by uranyl-carbonato to one dominated by uranyl-calcium-carbonato complexes correlates with the observed decrease in U(VI) reduction by \( S. \text{ putrefaciens} \) (Figure 3). This is reflected in the biogeochemical model by \( k_1 \) and \( k_{1b} \) having more than an order of magnitude difference in value: \( k_1 = 7.5 \times 10^{-6} \text{s}^{-1}, k_{1b} = 3.1 \times 10^{-7} \text{s}^{-1} \) (ie \( k_1 >> k_{1b} \)). Reduction of the various uranyl species, when coupled with lactate oxidation, results in appreciable shifts in apparent energy yield but does not systematically decrease with the ternary Ca complexes (Table 1).

Uranium(VI) Reduction

Reduction of U(VI) by \( S. \text{ putrefaciens} \) transpires at all Ca concentrations and with all 3 types of ferric (hydr)oxides investigated; however, both the rate and the extent of U(VI) reduction are dependent on Ca concentration – with an inverse relation between reduction rate and Ca concentration (Figure 3). With ferrihydrite, \( U_{(aq)} \) is reduced to 24% of its original concentration
of 0.168 mM after 528 h when no Ca is present in solution. In contrast, addition of Ca (initial concentrations of 0.4, 0.6, and 0.8 mM) results in a 50% reduction in $U_{(aq)}$ after 528 h. In the case of goethite, $U_{(aq)}$ is reduced to 2% of starting concentration after 528 h with no Ca, while with Ca, reduction is diminished and thus 20 to 35% of the initial U (0.168 mM) remains in solution after 528 h. With hematite, reduction ranges from 100% (no Ca) to 80% depending on the Ca concentration (Figure 3). The addition of Ca to the no-Fe control system has the most dramatic affect on U(VI) reduction. After 528 h of reaction, the extent of U reduction decreases from 90% to 55% of the initial concentration with the addition of Ca — consistent with the findings of Brooks et al.(2003).

**Iron(III) Reduction**

The concentration of Fe in the aqueous phase increased during the reaction period for ferrihydrite at all Ca concentrations (Figure 4), while the concentration of U in the aqueous phase decreased. The concentration of Fe generated after 528 h ranges from 0.11 mM in the absence of Ca, a concentration and rate comparable to that reported previously (Wielinga et al., 2000), to 0.29 mM in the presence of 0.8 mM Ca. Dissolved Ca appears to correlate with the concentration of Fe(II) produced, supporting the premise that Ca renders U(VI) less favorable as a terminal electron acceptor, thus making Fe(III) the dominant electron acceptor. In contrast to the ferrihydrite system, negligible amounts (<0.2 $\mu$mol L$^{-1}$) of aqueous Fe are detected in goethite or hematite systems (and in the no-Fe controls) at any Ca concentration; minimal Fe reduction transpired (less than the re-adsorption capacity of Fe(II) on goethite or hematite). The differences observed among the Fe (hydr)oxides are consistent with the surface area and free
energy yields, where Fe is, at least initially, more available for bioreduction when present as ferrihydrite than either goethite or hematite (Hansel et al., 2003, 2004).

In order to quantify the impact of Fe(III) on U(VI) reduction used for model input, the concentration of Fe(II) produced over 100 h of incubation in ferrihydrite systems was measured. The concentration of total Fe(II) (sum of aqueous and adsorbed components) increased during the reaction period at all Ca concentrations and reached a value of $0.16 \pm 0.01$ mM for all Ca concentrations (including no Ca) after 100 h of reaction. This concentration and rate are comparable to previous studies (Wielinga et al., 2000). Above, we report an increase in $\text{Fe}^{(aq)}$ for 0.8 mM Ca compared to 0 mM Ca but this did not appear until $>500$ h (Stewart et al., 2007).

**Time-Dependence of U(VI) Reduction.**

Uranium is removed from solution via reduction by *S. putrefaciens* with subsequent precipitation of biogenic $\text{UO}_2$ (s) in all systems studied but at varying rates. The rate of U removed from solution (expressed as the fraction of initial U) conforms to a first-order rate expression (Figure 5). Pseudo first-order rate constants, $k$ (h$^{-1}$), are larger for systems without than those with Ca (Table 4), with the variation being most pronounced for the iron (hydr)oxide-free system where the rate constant decreases from $13.7 \pm 0.3$ ($x10^{-3}$) h$^{-1}$ in the absence of Ca to $0.8 \pm 0.0$ ($x10^{-3}$) h$^{-1}$ when 0.8 mM Ca is added to the system. It follows that the rate constant in the presence of ferrihydrite changes the least, ranging from $4.1 \pm 0.2$ ($x10^{-3}$) h$^{-1}$ for 0 mM Ca to $2.4 \pm 0.2$ ($x10^{-3}$) h$^{-1}$ for 0.8 mM Ca. Pseudo first-order rate constants (Table 2) are linearly correlated to calcium concentrations between 0 and 1 mM (Figure 5). The above mentioned rate constants derived from a simple kinetic analysis are consistent with the full biogeochemical model results.
Simulation Results

Simulations of no-ferrihydrite experiments show relatively good fits to the measured data suggesting that the conceptual model used for, and particularly the rate dependence of, U(VI) reduction on aqueous UO$_2^{2+}$ speciation (Equation 9) interpret the data well (Figure 6). In some cases, particularly at later times (i.e. >70 h) for the 0 mM Ca and 0.4 mM Ca experiments, the simulation over predicts U(VI) reduction. This indicates that the dependence of U(VI) reduction on the no-Ca UO$_2^{2+}$ species is not precisely described by the first-order expression shown in Equation 9. Additional simulations (not shown) indicate that the model is sensitive to the exponent $x_1$, with a change from 1 to 1.05 making a noticeable difference in the results (i.e. up to 0.025 mM). The sensitivity to this parameter highlights the rate dependence on the concentration of UO$_2^{2+}$ species. However, we have used the same rate expressions and parameters for each simulation to highlight the effects of the change in Ca concentrations in the no-ferrihydrite experiments, and the competition between Fe(III) reduction and U(VI) reduction in the with-ferrihydrite experiments. As a result, slight differences in the rates between the experiments will not be captured by the simulations.

At early times in the no-ferrihydrite experiments (<10 h) the experimental results show a sharp decline in the U$_{(aq)}$ concentrations followed by a slight rebound that is not represented by the simulations. This behavior is likely attributed to rapid microbial reduction, followed by re-oxidation, before aqueous uranium begins to decrease at a more constant rate than is captured by the model.
In general, simulations of the with-ferrihydrite experiments predict the experimental data reasonably well, with the slight exception of the 0.8 mM Ca experiment (Figure 7). The good agreement between the simulated and measured data suggests that the conceptual model captures the important features of the geochemical system.

Discussion

Influence of Ca on U(VI) Reduction

There are several possible mechanisms by which Ca may diminish uranium reduction. First, Ca may render the reduction of uranyl less thermodynamically favorable. Second, Ca may kinetically limit reduction by yielding a dominant complex that either has poor-orbital overlap with the uranium reductase or hinders site accessibility. Finally, Ca could have a toxic affect on the microorganisms. Previous studies have eliminated the possibility of toxic effects of Ca or Ca-U-CO₃ to the bacteria (Brooks et al., 2003). Thermodynamic considerations illustrate that there is less than 10% difference in energy gained between the reduction of uranyl-carbonato and uranyl-calcium-carbonato species (Table 3). The Gibb’s free energy of reaction ($\Delta G_{\text{rxn}}$) for reduction of $(\text{UO}_2)_3\text{CO}_3(\text{OH})_3^-$ (-157.3 kJ mol⁻¹), the dominant aqueous species in the absence of calcium, and $\text{Ca}_2\text{UO}_2(\text{CO}_3)_3$ (-144.0 kJ mol⁻¹), the dominant species in the presence of calcium, are only 8% different—and both are highly favorable. Moreover, even with a Ca concentrations of 1 mM, the free energy of reaction for reduction of U(VI) is comparable to the energy yield from the reduction of ferrihydrite (under the conditions of this study). This gives rise to the hypothesis that the reduction of U(VI) by *S. putrefaciens* may have an unfavorable orbital symmetry, be sterically hindered, or have a high activation energy associated with uranyl-calcium-carbonato complex dissociation. This phenomenon appears to extends to chemical
reduction of U(VI) by Fe(II) in the presence of the Ca-U-CO$_3$ complexes as noted by Nico et al. (2009) and Stewart et al. (2009)

**Rate Determining Steps in U(VI) Reduction**

Although a multitude of U(VI) species reside under the conditions evaluated within this study, there are two dominant species, Ca$_2$UO$_2$(CO$_3$)$_3$ and (UO$_2$)$_2$CO$_3$(OH)$_3$ $^-$, under the geochemical conditions of this study; a third species, UO$_2$(CO$_3$)$_2$$^{2-}$, resides at appreciable fractions of the total uranium. This allows for a simplified kinetic analysis as compared to the one above (Model Development Section). Considering these three species, reduction reactions coupled with lactate oxidation (Table 1) are described by the stoichiometries of reactions 1 to 3.

\[
\begin{align*}
C_3H_5O_3^- + (UO_2)_2CO_3(OH)_3^- & \rightarrow C_2H_3O_2^- + 2HCO_3^- + 2UO_2 + H_2O + H^+ \\
\frac{1}{2}C_3H_5O_3^- + UO_2(CO_3)_2^{2-} + H_2O & \rightarrow \frac{1}{2}C_2H_3O_2^- + \frac{5}{2}HCO_3^- + UO_2 + \frac{1}{2}H^+ \\
\frac{1}{2}C_3H_5O_3^- + Ca_2UO_2(CO_3)_3 + H_2O + \frac{1}{2}H^+ & \rightarrow \frac{1}{2}C_2H_3O_2^- + \frac{7}{2}HCO_3^- + UO_2 + 2Ca^{2+}
\end{align*}
\]

Given that the reactions are unidirectional, the pH was maintained constant, and U(VI) reduction proceeds independent of lactate (enzyme substrate) concentration under the conditions and reaction period studied (Figure 3), the operative rate determining, microbially-mediated, reduction reactions can be simply expressed:

\[
\begin{align*}
(UO_2)_x(CO_3)_y \overset{k_1}{\longrightarrow} UO_2(s) \hspace{3cm} (4)
\end{align*}
\]

\[
\begin{align*}
Ca_2UO_2(CO_3)_3 \overset{k_2}{\longrightarrow} UO_2(s) \hspace{3cm} (5)
\end{align*}
\]
where in reaction 4 $(\text{UO}_2)_{x}(\text{CO}_3)_y$ represents the combined contributions of the carbonato species $(\text{UO}_2)_2\text{CO}_3(\text{OH})_3^-$ and $\text{UO}_2(\text{CO}_3)_2^{2-}$, both of which are present in appreciable proportions under the reaction conditions examined.

Under conditions where the ternary calcium complex of U(VI) dominates the aqueous speciation, reduction may proceed either by reaction 5 or through the small proportion of carbonato complexes without calcium (reaction 4). In the latter case, inter-species conversion must then be described as well,

$$\text{Ca}_2\text{UO}_2(\text{CO}_3)_3 + \text{H}^+ \underset{k_{-3}}{\overset{k_3}{\rightleftharpoons}} \text{UO}_2(\text{CO}_3)_2^{2-} + \text{HCO}_3^- + 2\text{Ca}^{2+} \quad (6a)$$

Given the excess concentration of bicarbonate and calcium relative to uranium, and the constant pH, the inter-species conversion simplifies to:

$$\text{Ca}_2\text{UO}_2(\text{CO}_3)_3 \underset{k_{-3}}{\overset{k_3}{\rightleftharpoons}} \text{UO}_2(\text{CO}_3)_2^{2-} \quad (6b)$$

The reactions expressed in reactions 4 and 5 thus appear to represent the rate controlling processes for uranyl reduction, with inter-conversion of uranyl species potentially important within Ca containing solutions. On the basis of U(VI) reduction rates in the presence and absence of Ca, reaction 4 proceeds at a much faster rate than reaction 5 (i.e., $k_1 \gg k_2$). With the selective removal of the carbonato complexes, reaction 6 would proceed in the forward direction during reduction. Reaction 4 thus expresses the rate controlling reaction for non-Ca bearing solutions, while reaction 5 and the coupling of reactions 4 and 6 would operate in series (reaction 7) as the rate influencing reactions in Ca-bearing solutions.
This correlates well with simulation results that indicate the reduction rate constant for uranyl-carbonato species, \(k_1 = 7.5 \times 10^{-6} \text{s}^{-1}\), is more than 10 times the rate constant for uranyl-calcium-carbonato complexes, \(k_{1b} = 3.1 \times 10^{-7} \text{s}^{-1}\), suggesting that the rate of reduction of uranyl-calcium-carbonato complexes is appreciably lower but not zero.

**Impact of Fe(III) (hydr)oxides on U(VI) Reduction**

Ferric (hydr)oxides impart an interesting complexity on uranium reduction. On the one hand, they can serve as competing electron acceptors of U(VI) and as oxidants of U(IV) (under specific reaction conditions) (Figure 1). On the other hand, they can also regulate the dissolved concentration of Ca, decreasing the proportion of the Ca-U ternary complex, and enhancing U(VI) reduction.

Ferrihydrite has the highest redox potential of the Fe(III) phases investigated here, and indeed it is the only Fe(III) (hydr)oxide to decrease the extent of U(VI) reduction. As noted above, ferrihydrite may either compete as an electron acceptor in microbial respiration or may act as an oxidant of biogenic UO₂. A series of recent reports indicate reoxidation of uraninite by Fe(III) at high concentrations of HCO₃⁻ resulting from bacterial respiration (Sani et al., 2005; Senko et al., 2005; Wan, 2005). However, under the conditions of our study (HCO₃⁻ < 6 mM, 0.168 U(VI), and pH 7), even the oxidation of biogenic UO₂ to Ca₂UO₂(CO₃)₃ by ferrihydrite, the most viable reaction, would be thermodynamically favorable only at Fe(II) concentrations less than 0.025 mM Fe(II) during the initial stages of reduction and 0.050 mM at late stages. These Fe(II) values
are exceeded during the initial stages of reaction ($t < 100$ h) and reoxidation of U(IV) therefore does not appear operative in this system.

Ferrihydrite, however, is not a static phase during Fe(II) production and instead undergoes a cascade of secondary reactions, dominantly leading to goethite or magnetite depending on the Fe(II) concentrations (Hansel et al., 2004). With progressive incubation, the ferrihydrite system responds in a similar way to goethite and hematite, consistent with a shift in mineralogy from ferrihydrite to goethite during dissimilatory iron reduction (Hansel et al., 2003, 2004). Hansel et al. (2003) report a threshold limit of 0.4 mmol Fe(II)/g-ferrihydrite below which ferrihydrite converts via Oswald ripening primarily to goethite/lepidocrocite and above which it converts to both goethite and magnetite. Iron(II) levels produced in the ferrihydrite system do not exceed this threshold when Ca is present until 171 h (0.52 mM, 1.3 mmol Fe(II)/g-ferrihydrite), and would therefore lead to pronounced generation of goethite.

In addition to serving as a competing electron acceptor, Fe(III) (hydr)oxides influence U(VI) reduction through adsorption of Ca, diminishing its aqueous concentrations and thereby decreasing the proportion of the apparently less reactive $\text{Ca}_2\text{UO}_2(\text{CO}_3)_3$ species. The impact is pronounced with goethite and hematite (Figure 3). Thus, the inhibitory effect of Ca on U(VI) reduction may be diminished when competing sinks for Ca are present; ultimately, the dissolved concentration of Ca in equilibrium with the uranyl species is the controlling factor. Finally, all of the Fe(III) (hydr)oxides, and ferrihydrite in particular, have an intriguing impact on U(VI) reduction in the presence of Ca. While ferrihydrite acts as a competing electron acceptor for microbial reduction of U(VI), as Ca concentrations increase, U(VI) reduction is promoted (rather
than retarded) in systems with ferrihydrite as compared to those without. Under the specific geochemical conditions in this study, ferrihydrite does not impact the rate or extent of uranium reduction during the first 100 h of incubation without Ca. In systems containing Ca, however, ferrihydrite appears to increase rates of U(VI) reduction compared to systems without ferrihydrite; at Ca concentrations > 0.4 mM, however, uranium reduction rate becomes independent of Ca concentration. This gives rise to two hypotheses on ferrihydrite’s influence. One possibility is that Ca is adsorbing to the surface of ferrihydrite, offsetting equilibrium with the uranyl-calcium-carbonato complexes. However, aqueous measurements indicate that Ca concentrations are identical in systems with and without ferrihydrite. A second, more substantiative explanation is that ferrihydrite adsorbs both free-uranyl and uranyl-carbonato complexes that are subject to microbial or Fe(II)-induced (Liger et al., 1999) reduction. In this scenario, Ca (aq) competes with the surface of ferrihydrite for uranyl and uranyl-carbonato species, with the Ca complexed uranium having a diminished reduction rate compared to free or adsorbed uranium. More information on changes in uranyl speciation in the presence of solid phases is provided in Stewart et., (2010).

**Implications for Uranium Reduction and Mobility**

Despite the observation of uranium(VI) reduction and subsequent precipitation of UO₂, both the rate and extent of reduction in complex geochemical settings are potentially diminished by aqueous and solid phase constituents of soils and sediments. The presence of calcium in carbonate-bearing solutions promotes the formation of ternary complexes, Ca₂UO₂(CO₃)₃ and CaUO₂(CO₃)₃²⁻, which have a profound impact on U biogeochemistry by decreasing both microbial and abiotic reduction rates. Iron (hydr)oxides, in contrast, have non-linear effects on
U(VI) reduction. Goethite and hematite act as sorbents of Ca, and as a result decrease the proportion of the less reducible Ca$_2$UO$_2$(CO$_3$)$_3$ species. At higher calcium concentrations, ferrihydrite has the ability to mitigate speciation changes invoked by the predominance of these complexes, thus increasing the rate of microbial reduction of uranium compared to systems with no ferrihydrite. Ultimately, U(VI) speciation, induced by both aqueous constituents - particularly the formation of Ca-UO$_2$-CO$_3$ complexes and equilibrium solid phases, the presence of competing terminal electron acceptors, and potential oxidants, will control the rate and extent of reduction.

**Acknowledgments**

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References


### Tables

Table 1. Redox reactions for various U(VI) and Fe(III) species coupled with lactate oxidation (ferrihydrite, Fe(OH)$_3$; goethite, FeOOH; hematite, Fe$_2$O$_3$).

<table>
<thead>
<tr>
<th>Redox Reaction</th>
<th>$\Delta G_{\text{reaction}}, \text{kJ/mol}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + 4\text{Fe(OH)}_3 + 7\text{H}^+ \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 4\text{Fe}^{2+} + \text{HCO}_3^- + 10\text{H}_2\text{O}$</td>
<td>$-291.1^{(1)}$</td>
</tr>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + 4\text{FeOOH} + 7\text{H}^+ \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 4\text{Fe}^{2+} + \text{HCO}_3^- + 6\text{H}_2\text{O}$</td>
<td>$-245.1^{(2)}$</td>
</tr>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + 2\text{Fe}_2\text{O}_3 + 7\text{H}^+ \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 4\text{Fe}^{2+} + \text{HCO}_3^- + 4\text{H}_2\text{O}$</td>
<td>$-210.1^{(2)}$</td>
</tr>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + 2\text{UO}_2(\text{CO}_3)_3^{4-} + 2\text{H}_2\text{O} + \text{H}^+ \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 7\text{HCO}_3^- + 2\text{UO}_2$</td>
<td>$-155.1^{(3)}$</td>
</tr>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + 2\text{UO}_2(\text{CO}_3)_2^{2-} + 2\text{H}_2\text{O} \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 5\text{HCO}_3^- + 2\text{UO}_2 + \text{H}^+$</td>
<td>$-96.9^{(3)}$</td>
</tr>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + (\text{UO}_2)_2\text{CO}_3(\text{OH})_3^- \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 2\text{HCO}_3^- + 2\text{UO}_2 + \text{H}_2\text{O} + \text{H}^+$</td>
<td>$-114.5^{(3)}$</td>
</tr>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + 2\text{Ca}_2\text{UO}_2(\text{CO}_3)_3^{3+} + 2\text{H}_2\text{O} + \text{H}^+ \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 7\text{HCO}_3^- + 2\text{UO}_2 + 4\text{Ca}^{2+}$</td>
<td>$-55.9^{(4)}$</td>
</tr>
<tr>
<td>$\text{C}_3\text{H}_5\text{O}_3^- + 2\text{CaUO}_2(\text{CO}_3)_2^{2-} + 2\text{H}_2\text{O} + \text{H}^+ \rightarrow \text{C}_2\text{H}_3\text{O}_2^- + 7\text{HCO}_3^- + 2\text{UO}_2 + 2\text{Ca}^{2+}$</td>
<td>$-94.3^{(5)}$</td>
</tr>
<tr>
<td>$2\text{Fe(OH)}_3 + 3\text{H}^+ + 3\text{HCO}_3^- + \text{UO}_2 + 2\text{Ca}^{2+} \rightarrow 2\text{Fe}^{2+} + 6\text{H}_2\text{O} + \text{Ca}_2\text{UO}_2(\text{CO}_3)_3$</td>
<td>$-117.6^{(1,4)}$</td>
</tr>
<tr>
<td>$2\text{Fe(OH)}_3 + 4\text{H}^+ + \text{UO}_2 + 0.5\text{HCO}_3^- \rightarrow 2\text{Fe}^{2+} + 4.5\text{H}_2\text{O} + 0.5(\text{UO}_2)_2\text{CO}_3(\text{OH})_3^-$</td>
<td>$-88.3^{(1,3)}$</td>
</tr>
<tr>
<td>$2\text{FeOOH} + \text{UO}_2 + 3\text{H}^+ + 3\text{HCO}_3^- + 2\text{Ca}^{2+} \rightarrow \text{Ca}_2\text{UO}_2(\text{CO}_3)_3^{3+} + 2\text{Fe}^{2+} + 4\text{H}_2\text{O}$</td>
<td>$-95.6^{(2,4)}$</td>
</tr>
<tr>
<td>$2\text{FeOOH} + \text{UO}_2 + 4\text{H}^+ + 0.5\text{HCO}_3^- \rightarrow 0.5(\text{UO}_2)_2\text{CO}_3(\text{OH})_3^+ + 2\text{Fe}^{2+} + 2.5\text{H}_2\text{O}$</td>
<td>$-65.3^{(2,3)}$</td>
</tr>
<tr>
<td>$\text{Fe}_2\text{O}_3 + \text{UO}_2 + 3\text{H}^+ + 3\text{HCO}_3^- + 2\text{Ca}^{2+} \rightarrow \text{Ca}_2\text{UO}_2(\text{CO}_3)_3 + 2\text{Fe}^{2+} + 3\text{H}_2\text{O}$</td>
<td>$-77.1^{(2,4)}$</td>
</tr>
<tr>
<td>$\text{Fe}_2\text{O}_3 + \text{UO}_2 + 4\text{H}^+ + 0.5\text{HCO}_3^- \rightarrow 0.5(\text{UO}_2)_2\text{CO}_3(\text{OH})_3^+ + 2\text{Fe}^{2+} + 1.5\text{H}_2\text{O}$</td>
<td>$-47.8^{(2,3)}$</td>
</tr>
</tbody>
</table>

1) Majzlan et al. (2004)  
3) Guillaumont et al. (2003)  
4) Bernhard et al. (2001)  
5) Dong et al. (2006)
Table 2. Pseudo-first order rate coefficients ($k_{obs}$) at various calcium concentrations for the different Fe(III) (hydr)oxide systems.

<table>
<thead>
<tr>
<th>Ca (mM)</th>
<th>Ferrihydrite</th>
<th>Goethite</th>
<th>Hematite</th>
<th>Iron oxide-free</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ln $k_{obs}$ = -0.650(Ca) - 5.56</td>
<td>Ln $k_{obs}$ = -1.72(Ca) - 4.55</td>
<td>Ln $k_{obs}$ = -1.46(Ca) - 4.72</td>
<td>Ln $k_{obs}$ = -3.57(Ca) - 4.46</td>
</tr>
<tr>
<td>0</td>
<td>4.1±0.2</td>
<td>11.9±0.6</td>
<td>9.7±0.4</td>
<td>13.7±0.3</td>
</tr>
<tr>
<td>0.4</td>
<td>2.6±0.0</td>
<td>4.6±0.0</td>
<td>4.1±0.1</td>
<td>2.0±0.1</td>
</tr>
<tr>
<td>0.6</td>
<td>2.7±0.0</td>
<td>3.1±0.0</td>
<td>4.0±0.1</td>
<td>1.3±0.1</td>
</tr>
<tr>
<td>0.8</td>
<td>2.4±0.2</td>
<td>3.3±0.5</td>
<td>2.0±0.1</td>
<td>0.8±0.0</td>
</tr>
</tbody>
</table>

*Values having an associated error of 0.0 as the composite error are reported relative to the mean and these particular data points lie on the mean value.*
Table 3. Gibb’s free energy of reaction ($\Delta G_{\text{reaction}}$) for Fe(III) and U(VI) reduction coupled with lactate oxidation at varying Ca(aq) concentrations. Values are calculated for dominant aqueous species as indicated in Table 2 and are adjusted for experimental conditions (pH=7, [CO$_3$]$_T$=3.8mM, [U(VI)] = 0.168mM).

<table>
<thead>
<tr>
<th></th>
<th>$\Delta G_{\text{reaction}}$ (KJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 mM Ca$^{2+}$</td>
</tr>
<tr>
<td>Fe(OH)$_3$/lactate</td>
<td>-139.5</td>
</tr>
<tr>
<td>Fe$_2$O$_3$/lactate</td>
<td>-58.27</td>
</tr>
<tr>
<td>Ca$_2$UO$_2$(CO$_3$)$_3$/lactate</td>
<td>-148.3</td>
</tr>
<tr>
<td>UO$_2$(CO$_3$)$_2^{2-}$/lactate</td>
<td>-158.8</td>
</tr>
<tr>
<td>UO$_2$(CO$_3$)$_3^{4-}$/lactate</td>
<td>42.91</td>
</tr>
<tr>
<td>(UO$_2$)$_2$CO$_3$(OH)$_3$/lactate</td>
<td>-158.60</td>
</tr>
</tbody>
</table>
**Figure Captions**

Figure 1. U(VI)/U(IV) and Fe(III)/Fe(II) redox couples at pH=7, [CO$_3$]$_T$=3 mM, and [Ca]=1mM illustrating the overlapping nature of U and Fe redox couples.

Figure 2. Fractional species of U(VI) as a function of Ca concentration (mM) depicted as the sum of U-CO$_3$ and U-Ca-CO$_3$ species (pH = 7, [CO$_3$]$_T$ = 3.8 mM).

Figure 3. Temporal changes in uranyl concentration induced by *S. putrefaciens* at Ca concentrations ranging from 0 to 0.8 mM in the presence of a) ferrihydrite, b) goethite, c) hematite and d) Fe -free system.

Figure 4. Temporal changes in Fe (aq) concentration induced by *S. putrefaciens* in the presence of ferrihydrite.

Figure 5. Relation between observed pseudo first-order rate coefficient and aqueous Ca concentrations for ferrihydrite, goethite, hematite, and Fe (hydr)oxide-free systems.

Figure 6. Aqueous phase U (individual species and total U) during U(VI) reduction by *S. putrefaciens* with lactate as a function of time for systems containing (A) 0 mM Ca, (B) 0.4 mM Ca, (C) 0.6 mM Ca, and (D) 0.8 mM Ca. Experimental data are represented by squares and simulations by lines.

Figure 7. Aqueous phase U and total Fe(II) produced during U(VI) and Fe(III) (ferrihydrite) reduction by *S. putrefaciens* with lactate as a function of time for systems containing (A) 0 mM Ca, (B) 0.4 mM Ca, (C) 0.6 mM Ca, and (D) 0.8 mM Ca. Experimental data are represented by squares and simulations by lines.
Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.
CaUO$_2$(CO$_3$)$_3$$^{-2}$

Ca$_2$UO$_2$(CO$_3$)$_3$(aq)

(UO$_2$)$_2$CO$_3$(OH)$_3$

UO$_2$CO$_3$(aq)

UO$_2$(CO$_3$)$_2$$^{2-}$

UO$_2$(CO$_3$)$_3$$^{4-}$

U(aq) Data

U(aq) Tot

Figure 6.
Figure 7.