The Influence of Calcium Carbonate Grain Coatings on Contaminant Reactivity in Vadose Zone Sediments

(Project Number: 70121)

Principal Investigator

John M. Zachara
Pacific Northwest National Laboratory
P.O. Box 999, MSIN K8-96
Richland, WA 99352
509-376-3254 (phone)
509-376-3650 (fax)
mailto:john.zachara@pnl.gov

Co-Investigators

Scott Chambers
Pacific Northwest National Laboratory
P.O. Box 999, MSIN K8-93
Richland, WA 99352
509-376-1766 (phone)
509-376-5106 (fax)
sa.chambers@pnl.gov

Gordon E. Brown, Jr.
Stanford University
Stanford, CA 94305
650-723-9168 (phone)
650-725-2199 (fax)
gordon@pangea.stanford.edu

Carrick M. Eggleston
University of Wyoming
P.O. Box 3006
Laramie, WY 82071
307-766-6769 (phone)
307-766-6679 (fax)
mailto:carrick@uwyo.edu
Research Objective

The primary objective of this project is to investigate the role of calcium carbonate grain coatings on adsorption and heterogeneous reduction reactions of key chemical and radioactive contaminants in sediments on the Hanford Site. Research will ascertain whether these coatings promote or discourage contaminant reaction with sediment mineral particles, and whether calcium carbonate phases resulting from waste-sediment reaction sequester contaminants through coprecipitation. The research will provide new conceptual models of contaminant reaction/retardation processes in Hanford sediments (for $^{90}\text{Sr}^{2+}$ and Cr(VI)O$_4^{2-}$ primarily) and improved geochemical models to forecast the future behavior of in-ground contaminants.

Research Progress and Implications

This project was initiated in fiscal year 1999 and is in its final year. Two primary research activities have been undertaken: 1) Characterization of $^{90}\text{Sr}^{2+}$ adsorption and exchange in pristine and contaminated calcareous Hanford sediments and 2) investigation of the role of calcium carbonate grain coatings on the heterogeneous reduction of Cr(VI)O$_4^{2-}$ by magnetite in model and sediment systems.

Characterization of Strontium Adsorption and Exchange

The ion exchange behavior of Sr$^{2+}$ was studied in pristine, calcareous sediments in both Na and Ca electrolytes. The sediment samples were drawn from Hanford’s B-BX-BY and T-TX-TY tank farms. Strontium adsorption in both systems displayed strong dependence on electrolyte concentration, consistent with an ion exchange process. The sand-textured sediments exhibit a small cation exchange capacity (approximately $10^{-5}$ eq/g) resulting from the presence of detrital layer silicates. A multicomponent cation exchange model (Na-Ca-Sr) was developed for the entire data set that includes three ion exchange selectivity coefficients $K_{\text{Na-Sr}}$, $K_{\text{Ca-Sr}}$, and $K_{\text{Na-Ca}}$ (where $K$ = the Vanselow selectivity coefficient) and the measured cation exchange capacities. Because the Hanford sediments contain appreciable ion-exchangeable $^{86/87}\text{Sr}^{2+}$, the adsorption process of contaminant $^{90}\text{Sr}^{2+}$ (which is typically present at 3 to 4 orders of magnitude lower concentration than indigenous $^{86/87}\text{Sr}^{2+}$) is one of isotopic exchange with the native Sr pool. The $^{90}\text{Sr}^{2+}$-$K_d$ is therefore controlled by the $K_d$ of the indigenous Sr pool, which, in turn, is predictable from our developed multicomponent cation exchange model. Adsorption experiments with $^{86/87/90}\text{Sr}^{2+}$ are currently being performed to document the predictability of these relationships with water compositions characteristic of different Hanford waste-release scenarios.

In other experiments, we have studied the mineralogic association of sorbed $^{90}\text{Sr}^{2+}$ and its desorption kinetics from contaminated sediments collected beneath leaked tank B-110 in Hanford’s B tank farm complex. The waste stream leaked to these sediments was from the
Sr-recovery process. We have succeeded in isolating $^{90}$Sr$^{2+}$-containing particles from the sediments (Figure 1) and now are characterizing these in detail using various forms of electron microscopy. We have contacted one sediment sample from the B-110 borehole that contained approximately 10,000 pCi/g of sorbed $^{90}$Sr$^{2+}$ with various electrolytes intended to desorb

- ion-exchanged $^{90}$Sr$^{2+}$ (0.01, 1, and 5 mol/L NaNO$_3$; 0.05 mol/L Ca(NO$_3$)$_2$)
- coprecipitated $^{90}$Sr$^{2+}$ (1 mol/L NaOAc at pH 5; 0.5 mol/L HCl) (Figure 2).

Figure 1. Samples of B-110 sediment were mounted on TEM stubs and imaged by phosphor-luminescence. The dark areas are mineral grains or aggregate containing $^{90}$Sr. These grains were imaged by scanning electron microscopy and then were selectively removed for detailed mineralogic characterization and spatial Sr analysis.

![Phosphor-luminescence image of B-110 sediment](image)

Figure 2. Time-dependent desorption of $^{90}$Sr from contaminated B-110 sediment. $^{90}$Sr was released to the sediment over 30 years ago in the form of a Sr recovery waste containing Na-HCO$_3$ and complexants. Extraction with 0.01, 1, and 5 mol/L Na and 0.05 mol/L (Ca(NO$_3$)$_2$ removes inexchangeable $^{90}$Sr, while NaOAc at pH 5 and 0.5 mol/L HCl dissolve host carbonate phases as well.
The experiments have demonstrated that approximately 75% of the sorbed $^{90}\text{Sr}^{2+}$ is associated with the exchange complex, and 25% is associated with carbonate grain coatings that have resulted from waste-sediment reaction. These data now are being modeled with our multicomponent exchange model linked with a precipitation model. Collectively, these results provide a basis for predicting the geochemical behavior of $^{90}\text{Sr}^{2+}$ in various waste streams and subsurface sediments at the Hanford site.

**Investigation of Calcium Carbonate Grain Coatings**

The growth of complete calcium carbonate overlayers on synthetic magnetite (001) surfaces was accomplished by a biomineralization approach termed the polymer-induced liquid precursor (PILP) method. During this solution-based approach, poly-aspartate coats the magnetite surface and acts as a surfactant upon which primarily vaterite nucleates as a continuous film. An atomic force microscopy micrograph taken during early growth stages displayed a beta-sheet-like layer, reminiscent of poly-aspartate, with a carbonate overlayer nucleating atop. These calcium carbonate overlayers of vaterite crystal structure interfere with the reduction of $\text{Cr(VI)}\text{O}_4^{2-}$ at the magnetite-water interface. Films as thin as 20 to 25 angstroms inhibit the reduction of aqueous chromate throughout the pH range of 6 to 8 for exposures less than one hour (Figure 3). Depending upon the carbonate film thickness, however, $\text{Cr(VI)}\text{O}_4^{2-}$ reduction did occur after

![Figure 3](image.png)

**Figure 3.** High-energy resolution Cr 2p core-level spectra for synthetic magnetite(001) covered with calcium carbonate (gray) and not covered (black). Reference spectra for Cr(VI) and Cr(III) also are shown.
extended reaction times due to dissolution of the calcium carbonate coatings. Surprisingly,
however, secondary ion mass spectrometry (SIMS) analysis of samples from extended reaction
showed that Cr was concentrated on the magnetite surface in areas of high carbon and less so in
areas of high Fe content. This was puzzling until the experiment was performed in which the
Cr(VI)O$_4^{2-}$ solution was also saturated with calcium carbonate. Expecting a lower dissolution
rate of the coating due to a saturated solution and less Cr(VI)O$_4^{2-}$ reduction, we found, in
contrast, that there was a significant Cr(III) photoemission signature. This was due to
photoreduction of Cr(VI) to Cr(III) in the presence of adventitious carbon during analysis in the
x-ray photoemission chamber.

These findings provide a clear explanation for why Cr(VI) and Tc(VII) are stable valence states
in the Hanford subsurface, in spite of the presence of significant magnetite and ilmenite in the
sediments. Calcium carbonate grain coatings prevent heterogeneous reduction.

**Planned Activities**

Being well on in the last funded year of the project, most of our remaining activity is focused on
publication of results. Some lingering experimentation with the $^{90}\text{Sr}^{2+}$-containing
B-110 sediments and magnetite isolated from the Hanford sediment will continue to completion
in the remaining months of fiscal year 2002. Some of our research results from the
B-110 sediments will be incorporated into a Corrective Action Assessment Report for the
B-BX-BY Tank Farms in July, thereby providing key scientific support for an important decision
milestone at the Hanford Site.

**Information Access**

Droubay T and SA Chambers. 2001. Surface-sensitive Fe 2p photoemission spectra for alpha-
Fe$_2$O$_3$(0001): The influence of symmetry and crystal-field strength - Art. No. 205414. *Physical
Review B* 6420(20):5414-+


Zachara JM, P Lichtner, and SC Smith. 2002. Ion exchange of Sr$^{2+}$ in calcareous subsurface
sediment and isotopic exchange of $^{90}\text{Sr}^{2+}$ as a retardation mechanism. *Geochemica et

Zachara JM, P Lichtner, and SC Smith. 2002. Mineralogic residence and desorption kinetics
$^{90}\text{Sr}^{2+}$ from contaminated subsurface sediments. *Environmental Science and Technology.*
Submitted.