Old Dominion University

ODU Digital Commons

Chemistry & Biochemistry Faculty Publications

Chemistry & Biochemistry

2016

Plutonium Immobilization and Mobilization by Soil Organic Matter

Peter H. Santschi

Kathleen A. Schwehr

Chen Xu

Matthew Athon

Yi-Fang Ho

See next page for additional authors

Follow this and additional works at: https://digitalcommons.odu.edu/chemistry_fac_pubs



Part of the Environmental Chemistry Commons, and the Radiochemistry Commons

Original Publication Citation

Santschi, P. H., Schwehr, K. A., Xu, C., Athon, M., Ho, Y.-F., Hatcher, P. G., Didonato, N., & Kaplan, D. I. (2016). Plutonium immobilization and mobilization by soil organic matter (No. DOE-TXAM-06820). https://doi.org/10.2172/1240745

This Report is brought to you for free and open access by the Chemistry & Biochemistry at ODU Digital Commons. It has been accepted for inclusion in Chemistry & Biochemistry Faculty Publications by an authorized administrator of ODU Digital Commons. For more information, please contact digitalcommons@odu.edu.

Authors Peter H. Santschi, Kathleen A. Schwehr, Chen Xu, Matthew Athon, Yi-Fang Ho, Patrick G. Hatcher, Nic Didonato, and Daniel I. Kaplan	
	col

Plutonium Immobilization and Mobilization by Soil Organic Matter

¹PETER H. SANTSCHI, KATHLEEN A. SCHWEHR, CHEN XU, MATTHEW ATHON, YI-FANG HO,

Background: The human and environmental risks associated with Pu disposal, remediation, and nuclear accidents scenarios stems mainly from the very long half-lives of several of its isotopes. The SRS, holding one-third of the nation's Pu inventory, has a long-term stewardship commitment to investigation of Pu behavior in the groundwater and downgradient vast wetlands.

Pu is believed to be essentially immobile due to its low solubility and high particle reactivity to mineral phase or natural organic matter (NOM). For example, in sediments collected from a region of SRS, close to a wetland and a groundwater plume, ^{239,240}Pu concentrations suggest immobilization by NOM compounds, as Pu correlate with NOM contents. Micro-SXRF data indicate, however, that Pu does not correlate with Fe. However, previous studies reported Pu can be transported several kilometers in surface water systems, in the form of a colloidal organic matter carrier, through wind/water interactions. The role of NOM in both immobilizing or remobilizing Pu thus has been demonstrated. Our results indicate that more Pu (IV) than (V) was bound to soil colloidal organic matter (COM), amended at far-field concentrations. Contrary to expectations, the presence of NOM in the F-Area soil did not enhance Pu fixation to the organicrich soil, when compared to the organic-poor soil or the mineral phase from the same soil source, due to the formation of COM-bound Pu. Most importantly, Pu uptake by organic-rich soil decreased with increasing pH because more NOM in the colloidal size desorbed from the particulate fraction at elevated pH, resulting in greater amounts of Pu associated with the COM fraction. This is in contrast to previous observations with low-NOM sediments or minerals, which showed increased Pu uptake with increasing pH levels. This demonstrates that despite Pu immobilization by NOM, COM can convert Pu into a more mobile form.

Methods: We have determined ambient Pu concentrations and isotopic ratios with ICP-MS and alpha spectrometry, in samples collected from both SRS and Fukushima Prefecture [1-3]

Results: Sediment Pu concentrations in the SRS F-Area wetland were correlated to total organic carbon and total nitrogen contents and even more strongly to hydroxamate siderophore (HS) concentrations. The HS were detected in the particulate or colloidal phases of the sediments but not in the low molecular fractions (< 1000 Da). Macromolecules which scavenged the majority of the potentially mobile Pu were further separated from the bulk mobile organic matter fraction ("water extract") via isoelectric focusing experiment (IEF). An ESI FTICR-MS spectral comparison of the IEF extract and a siderophore standard (desferrioxamine; DFO) suggested the presence of HS functionalities in the IEF extract.

The cuticle material from Western wheatgrass (Agropythi Smithii), one of the dominant vegetation species at the RFETS, was extracted, and carbohydrates removed from the crude cuticle extract. Oxidation and depolymerization experiments of cutin, incoporation of carbohydrates, siderophore compounds, as well as the chelation of Fe(III) (as a surrogate of

²PATRICK G. HATCHER, NICOLE DIDONATO

³Daniel I. Kaplan

¹Department of Marine Sciences, Texas A&M University, Building 3029, Galveston,

TEXAS 77553 (*CORRESPONDENCE: SANTSCHI@TAMU.EDU)

²Department of Chemistry& Biochemistry, College of Sciences, Old Dominion University, Norfolk, VA 23529

³Savannah River National Laboratory, Aiken, SC 29808

Pu(IV)) during a simulated humification process, were assessed by ¹³C NMR, HRMAS, ATR-FTIR, etc.).

This study suggests that while HS are a very minor component in the sediment particulate/colloidal fractions, their concentrations greatly exceed those of ambient Pu, and HS may play an especially important role in Pu immobilization and re-mobilization in soils from different climatic regions, e.g., from grasslands in Colorado, wetlands in South Carolina, and paddy, urban, deciduous forest and coniferous forest soils from Japan.

References:

- 1. Xu, C. *et al.* Evidence for hydroxamate siderophores and other N-containing organic compounds controlling 239,240Pu immobilization and re-mobilization in a wetland sediment. *Environmental Science & Technology* **49**, 11458-11467, doi:10.1021/acs.est.5b02310 **(2015).**
- 2. Xu, C. *et al.* Plutonium Immobilization and Remobilization by Soil Mineral and Organic Matter in the Far-Field of the Savannah River Site, U.S. *Environmental Science & Technology* **48**, 3186-3195, doi:10.1021/es404951y **(2014).**
- 3. Xu, C. *et al.* Role of natural organic matter on iodine and 239,240Pu distribution and mobility in environmental samples from the northwestern Fukushima Prefecture, Japan. *Journal of Environmental Radioactivity* **153**, 156-166, doi:http://dx.doi.org/10.1016/j.jenvrad.2015.12.022 **(2016).**
- 4. Nicole DiDonato, Chen Xu, Peter H. Santschi, and Patrick G Hatcher. 2016. *A new perspective on structural components of Pu-mobilizing organic colloids from contaminated*, to be submitted to ES&T