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Reactive Transport of U and V from Abandoned Uranium Mine Wastes

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Abstract

The reactive transport of uranium (U) and vanadium(V) from abandoned mine wastes collected from the Blue Gap/Tachee Claim-28 mine site in Arizona was investigated by integrating flowthrough column experiments with reactive transport modeling, and electron microscopy. The mine wastes were sequentially reacted in flow-through columns at pH 7.9 (10 mM HCO₃⁻) and pH 3.4 (10 mM CH₃COOH) to evaluate the effect of environmentally relevant conditions encountered at Blue Gap/Tachee on the release of U and V. The reaction rate constants (k_m) for the dissolution of uranyl-vanadate (U-V) minerals predominant at Blue Gap/Tachee were obtained from simulations with the reactive transport software, PFLOTRAN. The estimated reaction rate constants were within 1 order of magnitude for pH 7.9 ($k_m = 4.8 \times 10^{-13} \text{ mol cm}^{-2} \text{ s}^{-1}$) and pH 3.4 ($k_m = 3.2 \times 10^{-13} \text{ mol cm}^{-2} \text{ s}^{-1}$) 10^{-13} mol cm⁻² s⁻¹). However, the estimated equilibrium constants (K_{eq}) for U–V bearing minerals were more than 6 orders of magnitude different for reaction at circumneutral pH (K_{ea} = $10^{-38.65}$) compared to acidic pH ($K_{eq} = 10^{-44.81}$). These results coupled with electron microscopy data suggest that the release of U and V is affected by water pH and the crystalline structure of U-V bearing minerals. The findings from this investigation have important implications for risk exposure assessment, remediation, and resource recovery of U and V in locations where U-Vbearing minerals are abundant.

Graphical abstract

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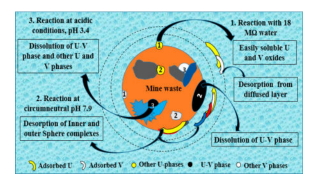
AUTHOR INFORMATION

Notes

The authors declare no competing financial interest.

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INTRODUCTION

The legacy of mill tailings and wastes from mining operations have resulted in the release of elevated concentrations of metals and radionuclides, which pose imminent environmental and human health concerns.^{1, 2} For instance, the negative health impacts of human exposure to uranium (U) and vanadium(V) through inhalation and ingestion have been well documented.^{3, 4} Uranium and V co-occur at numerous abandoned mine waste sites on the Navajo Nation territory near the four corners region of the southwestern U.S.^{5, 6} Uranium and other co-occurring metals from these mine wastes can be released into neighboring water resources affecting their water quality.

The aqueous and solid chemical speciation of U and V at mine waste sites could be affected by a variety of interfacial geochemical processes. These mine waste sites are usually exposed to ambient surface oxidizing conditions in which U(VI) is predominant.^{6, 7} Aqueous U(VI) can complex with OH^- and CO_3^{2-} , or react with alkaline earth metals and CO₃²⁻to form neutral or negatively charged ternary complexes (e.g., U-Ca-CO₃²⁻) that affect U mobility in water.^{8,9} Vanadium preferentially forms anionic $[V_{10}O_{26}(OH)_2^{4-},$ V₁₀O₂₇(OH)⁵⁻, V₁₀O₂₈⁶⁻, VO₂(OH)²⁻, VO₃(OH)²⁻, VO₄³⁻, V₂O₆(OH)³⁻, V₂O₇⁴⁻, $V_3O_9^{3-}$, and $V_4O_{12}^{4-}$], cationic [VO²⁺], and neutral [VO(OH)₃] species of oxyanions and aqueous complexes.^{10, 11} These oxyanions and aqueous complexes of V and U(VI) can adsorb onto charged surfaces of Al oxides, Fe oxides, and clay minerals.9, 11-18 Uranium and V can also be present in solid form as oxide, phosphate, carbonate, vanadate, and silicate mineral phases.^{19–22} For instance, their co-occurrence as carnotite $[K_2(UO_2)_2V_2O_8]$, tyuyamunite [Ca(UO₂)₂V₂O₈·(5–8)H₂O], metahewettite [CaV₆O₁₆·3(H₂O)], melanovanadite $[CaV_4O_{10}, 5(H_2O)]$, and V-bearing clays has been observed at the Rough Rock Sandstone of the Cretaceous Mesa Verde Group study region located in southwestern US.²³ Uranium oxides such as schoepite [(UO₂)₈O₂(OH)₁₂·12(H₂O)] are commonly occurring minerals that also influence uranyl solubility under oxic and suboxic environments between pH 6-9.24, 25 These geochemical processes, which are key to the aqueous and solid speciation of U and other co-occurring metals, should be considered when investigating the reactive transport of metals in abandoned mine wastes.

Reactive transport modeling can be a useful tool to investigate the interfacial mechanisms that affect the surface and subsurface mobilization of U and other metals. Previous studies have used reactive transport modeling to investigate the adsorption behavior of U(VI)

complexes onto alluvial aquifer sediments.^{26, 27} The subsurface desorption kinetics of U(VI) species suggest a nonequilibrium sorption behavior with Fe-oxides, Mn-oxides, and phyllosilicates.^{7, 28, 29} Uranium transport can also be attributed to dissolution of mineral phases. For example, U transport from contaminated subsurface Hanford sediments was affected by the dissolution of metatorbernite $[Cu(UO_2)_2(PO_4)_2 \cdot 8H_2O]$, cuprosklodowskite $(Cu[(UO_2)-(SiO_2OH)]_2 \cdot 6H_2O)$, and Na-boltwoodite $[NaUO_2(SiO_3OH) \cdot 1.5H_2O]$.^{28, 30, 31} The development of similar reactive transport models that represent environmentally relevant oxidizing conditions in abandoned mine waste sites is necessary to better understand the mobility of U, V, and other metals.

Previous laboratory-based studies have shown that U and V can interact through precipitation reactions forming uranyl vanadate minerals with potassium and calcium (e.g., carnotite [K₂(UO₂)₂(VO₄)₂·3H₂O] and tyuyamunite [Ca(UO₂)₂V₂O₈· 5–8(H₂O)]).^{21, 32} These studies used batch experiments and thermodynamic predictions to identify precipitation of uranyl vanadates as an effective method to immobilize U and V below the U-maximum contaminant level (MCL, 30 μ g L⁻¹) between pH 5.5–7.^{21, 32} The thermodynamic estimates in these studies were in agreement with those previously calculated by Langmuir.³³ However, these theoretically estimated thermodynamic constants are limited to synthetic minerals that have been subjected to specific reactivity conditions which could differ from those of U–V bearing minerals occurring in the environment.

A recent study conducted by our research group at the Blue Gap/Tachee Claim-28 mine site located in Northeastern Arizona reported that the dissolution of U–V bearing minerals affects the mobility of U and V at pH conditions observed in the field, which range from 3.8 to 7.4.⁶ Results showed that U and V co-occur in mine wastes predominantly as U(VI) and V(V). The Extended X-ray absorption fine structure (EXAFS) fittings suggest that U and V coordination in these mine waste samples is similar to that of minerals such as carnotite or tyuyamunite. Additionally, results from this study suggest dissolution of these U–V bearing minerals as a potential source to the elevated metal concentrations in neighboring surface water resources.⁶ Although this study represents a valuable first attempt to identify the co-occurrence of U and other metals at this abandoned mine waste site, the mechanisms affecting the reactive transport of U, V, and other metals remain unknown.

The objective of this study is to investigate the reactive transport of U and V from the Blue Gap/Tachee Claim-28 mine waste site through laboratory experiments, reactive transport modeling, and electron microscopy. In the current investigation we seek to determine the equilibrium and reaction rate constants for dissolution of these U–V bearing minerals at circumneutral and acidic pH, by modeling the batch and column experiments performed in this study with PFLO-TRAN.^{27, 28, 33} This study was developed based on our prior evidence that suggested U–V bearing mineral phases as the potential source to U and V release from Blue Gap/Tachee mine wastes.⁶ Our integrated approach of using microscopy and reactive transport modeling serves as a foundation to better understand the mechanisms affecting the reactivity of mine wastes and the transport of U and V under environmentally relevant conditions.

MATERIALS AND METHODS

Materials

We collected abandoned mine waste and background sediment samples from the Blue Gap/ Tachee Claim-28 mine site in January 2014 to identify the processes and phases that affect the U and V transport into neighboring surface and groundwater resources. Sediment samples from the site included mine waste and baseline reference soil (background soil) previously characterized by Blake et al, 2015.⁶ The Blake et al. study used powder X-ray diffraction to identify quartz (SiO₂; 59%) potassium feldspar (KAlSi₃O₈; 34%), and kaolinite (Al₂Si₂O₅(OH)₄; 7%) as the major mineralogical constituents. Additionally the cooccurrence of U and V, and predominance of U(VI) and V(V) were identified using microscopy, X-ray photoelectron spectroscopy (XPS), and X-ray absorption near edge spectroscopy (XANES) techniques. Furthermore, the coordination numbers obtained from EXAFS fits suggested the structure of the U–V bearing mineral to be similar to that of carnotite or tyuyamunite.⁶

Batch Experiments

We conducted batch experiments to model the U and V release at circumneutral (8.3) and acidic (3.8) pH, and determined the equilibrium and reaction rate constants for dissolution of U–V bearing mineral phases. The determined equilibrium constant was then used to obtain the reaction rate constants for U–V bearing mineral dissolution during column experiments. Briefly in the batch experiments, we reacted 0.1 g of mine wastes and background soil samples that were sieved to <63 μ m size (using U.S. standard sieve series number 230), with 50 mL 10 mM HCO₃⁻ (pH 8.3) and 10 mM C₆H₈O₆ (pH 3.8) through gentle shaking in 50 mL centrifuge tubes for 264 h, as described in Blake et al.⁶ During the experiments we collected 2 mL aliquots 0.5, 1, 2, 6, 24, 48, 96, and 264 h after the experiment began. We diluted these aliquots and analyzed for elemental concentrations using inductively coupled plasma optical emission spectroscopy and mass spectroscopy (ICP-OES/MS). All batch experiments were conducted in triplicate.

Continuous Flow through Column Experiments

We conducted column experiments to investigate the transport of U and V from Blue Gap/ Tachee sediments. An initial set of experiments consisted of reacting mine waste and background soil with 18M Ω water at pH 5.4 to evaluate the release of water-soluble U and V. Another set of experiments was performed in parallel for which the mine waste and background soil were sequentially reacted at pH 7.9 (using 10 mM HCO₃⁻) and pH 3.4 (using 10 mM CH₃COOH). These pH values and concentrations represent environmentally relevant conditions observed at the study site.⁶ We prepared samples by crushing and homogenizing sediments with a Spex shatter box and sieving through U.S. standard sieve with series 120-45 to obtain particle sizes between 125 and 355 μ m. The homogenized sieved fractions were then packed into cylindrical Pyrex glass columns with an inner diameter of 1 cm and a height of 10 cm using vacuum filter pumps. Solutions of 10 mM HCO₃⁻ and 10 mM CH₃COOH were prepared using 18M Ω deionized water, NaHCO₃ (powder), and 17.4N CH₃COOH. The prepared solutions were pumped against gravity through the packed columns at 0.9 mL min⁻¹ using a Masterflex L/S series 8-channel

peristaltic pump. The column experiments were conducted on eight columns in parallel (Figure S1). We chose 0.9 mL min⁻¹ based on existing literature^{7, 28, 34} as a typical groundwater rate to evaluate how solutions at circumneutral and acidic conditions would react with mine wastes. Columns 1, 2, 5, and 6 were packed with mine waste and columns 3, 4, 7, and 8 were packed with background soil. Columns 1-4 were reacted with 18MΩ water for 90 min with a total of 23 (columns 1-2) and 31 (columns 3-4) pore volumes (PV) at the rate of 3.53 and 2.65 cm³ PV⁻¹, respectively. Columns 5–8 were sequentially reacted for 1 week each (10 080 min) with 10 mM HCO₃⁻ and 10 mM CH₃COOH (Figure S1). The 1 week reaction time resulted in 2570 PV for columns 5 and 6, and 3450 PV for columns 7 and 8. Samples were collected at 10, 20, 30, 40, 50, 60, 70, 80, and 90 min of reaction time from the outlets of columns 1, 2, 3, and 4. Samples were collected from columns 5–8, at 30, 180, 540, 1440, 2880, 4320, 5760, 7,200 and 10 080 min after sequential reaction with 10 mM HCO₃⁻ and 10 mM CH₃COOH solutions. The sample aliquots were filtered using a $0.45 \,\mu m$ syringe filter and diluted accordingly for analyses. All experiments were performed at room temperature (25 °C) and, consequently, that temperature was set for the reactive transport of modeling.

Aqueous Chemical Analyses

We used a PerkinElmer Optima 5300DV inductively coupled plasma-optical emission spectrometer (ICP-OES) and a PerkinElmer NexION 300D (Dynamic Reaction Cell) inductively coupled plasma-mass spectrometer (ICP-MS) to obtain the trace metal concentrations in experimental aliquots and total acid extracts of mine waste and background soil. The solution pH was measured using a Thermo Scientific Orion Versastar Advanced Electrochemistry pH meter. Additional technical details of these methods are described in the Supporting Information.

Reactive Transport Modeling

We used PFLOTRAN a surface-subsurface multiphase, multicomponent reactive flow and transport model^{28, 29, 35} to determine the equilibrium constants for U–V bearing mineral dissolution at circumneutral and acidic pH (batch experiments) and identify the processes controlling the reactive transport of U and V from mine waste (column experiments).

To model the batch and column experiments we used, various U and V bearing aqueous species and mineral, their associated thermodynamic equilibrium constants (K_{eq}) and stoichiometric reaction coefficients (Tables S1 and S2), as inputs for PFLOTRAN. Most of the listed parameters were fixed, except the equilibrium constant of carnotite and reaction rate constants of the U–V bearing minerals, metaschoepite [UO₃·2H₂O] and rutherfordine [(UO₂)CO₃] which were varied to fit the experimental data and obtain the equilibrium and reaction rate constants for dissolution of U–V bearing minerals. At circumneutral pH, minerals such as metaschoepite and rutherfordine were considered in the model as they are commonly occurring oxides and carbonates of U that affect U transport.^{24, 36–38} In addition to metaschoepite and rutherfordine, surface complexation reactions for > SOUO₂OH and > SOHUO₂CO₃ were also considered in order to understand the effect of sorption on the reactive transport of U.³⁹ The equilibrium constants estimated for U–V bearing minerals by

modeling the batch experiments at circumneutral and acidic pH were used to estimate their reaction rate constants during column experiments.

Some key assumptions made in the model include the following: (1) U–V bearing minerals are the dominant U phase in the mine waste sample. (2) U–V bearing minerals have similar properties to those of carnotite. (3) U–V bearing mineral particles are spherical in shape. These assumptions were made on the basis of our previously published information on the Blue Gap/Tachee mine site, where the EXAFS fits of U–V bearing minerals (that were dominant in the mine waste) suggested similar coordination as carnotite or tyuyamunite. The spherical shape of the U–V bearing minerals was assumed for calculation simplifications.

Using PFLOTRAN, we modeled the water movement through the column considering fully saturated sediments under oxidizing conditions. The governing mass conservation equation that accounts for the unidirectional change in the total dissolved concentration of the *j*th primary species is given by (eq 1):

$$\frac{\partial}{\partial t}(\varphi \Psi_j) + \frac{\partial}{\partial z}(q\Psi_j - \varphi D\frac{\partial}{\partial z}\Psi_j) = Q_j - \sum_m v_{jm}I_m - \frac{\partial S_j}{\partial t}$$
(1)

where z [L] is the height of the column (10 cm), t [T] is the time for which column experiments were conducted ψ_j [ML⁻³] denotes the total concentration, D [L⁻² T⁻¹] denotes hydrodynamic dispersion coefficient, Q_j [M L⁻³ T⁻¹] is a source/sink term, I_m [M L⁻³ T⁻¹] is the reaction rate of the *m*th mineral, v_{jm} [–] is the stoichiometric coefficient for mineral reactions, S_j [ML⁻³] is the sorbed concentration, q [L T⁻¹] is Darcy's flux calculated to be approximately 1.15 cm min⁻¹ based on the flow rate used for the column experiments, 0.9 mL min⁻¹ and φ [–] is porosity. Dirichlet boundary conditions were used to model the 1D reactive transport of metals during reaction at pH 7.9 (using 10 mM HCO₃⁻) and pH 3.4 (using 10 mM CH₃COOH). Additional information and other supporting equations used in the model are explained in the Supporting Information. (SI eq 2–13)

Solids Characterization

Solid characterization using JEOL 2010 high resolution transmission electron microscopy (HR-TEM) was performed to determine the crystallinity of U–V bearing minerals in the mine waste sample. Synchrotron micro- X-ray fluorescence mapping (μ -SXRF) was conducted in beamline (BL) 10-2 in Stanford Synchrotron Radiation Lightsource (SSRL). Additional technical details of these methods are described in the Supporting Information.

RESULTS AND DISCUSSION

Mine Waste Reactivity in Columns Using 18MΩ Water (pH 5.4)

We investigated the potential release of water-soluble U and V from mine waste solids by reacting columns with 18M Ω water as a function of pore volumes and time (Figure S2). Aqueous concentrations as high as 2.3×10^{-7} M U and 74×10^{-6} M V were measured. For reference, the concentrations of U released from this reaction (~ 2.3×10^{-7} M) were similar to the concentrations observed in one of the water sources proximate to the site (spring

concentrations equal to 2.8×10^{-7} M), as was shown in a previous study conducted by our group. These mobilized concentrations were almost two times higher than the regulated U.S. Environmental Protection Agency maximum contaminant level (MCL) of U, that is, 1.3×10^{-7} M (30 μ g L⁻¹)⁴⁰. The U and V concentrations released after the reaction of background soil (control) with 18MΩ water were either below the MCL for U or below the detection limit of the ICP-OES/MS instruments. The presence of U and V in the effluent solution can be attributed to possible contributions from electrostatically bound chemical species in the diffuse layer or dissolution of soluble oxide phases, such as schoepite [(UO₂)₈O₂(OH)₁₂·12(H₂O)] and V oxides as observed in other investigations.^{41–43} The mobility of such labile aqueous species from mine waste is environmentally relevant. Further investigation of the reactive transport of U and V based on co-occurring U–V bearing mineral phases was conducted by sequentially reacting mine waste and background soil with circumneutral and acidic reagents.

Reactive Transport of U and V at Circumneutral pH

Batch Experiments at pH (8.3)—The reactive transport model developed for batch experiments reacting mine waste with 10 mM HCO₃⁻ identified dissolution of U-V bearing minerals as the key process affecting the dissolved concentrations of U and V (Figure 1A,B). The equilibrium constant (K_{eq}) for U–V bearing mineral dissolution was estimated to be 10^{-44.81} assuming the stoichiometry of carnotite (Table 1), which was almost 12 orders of magnitude lower than that reported in the literature for carnotite ($K_{eq} = 10^{-56.38}$).⁴⁴ The difference between the K_{eq} for dissolution of U–V bearing minerals and carnotite can be attributed to a difference in reactivity conditions and crystallinity between U–V bearing minerals in the mine wastes and carnotite. In addition, the estimated reaction rate constant $(k_{\rm m})$ for U–V mineral dissolution was 2.13×10^{-14} mol cm⁻² s⁻¹ (Table 1). Similarly, the estimated $k_{\rm m}$ values for precipitation of metaschoepite and rutherfordine were approximately 1×10^{-15} and 2×10^{-16} mol cm⁻² s⁻¹, respectively (Table S3). The nature of a reaction (dissolution or precipitation) can be identified from the sign convention of the reaction rate (Im) appearing in PFLOTRAN, that is, negative for dissolution and positive for precipitation. These results are in agreement with observations previously made at the nearby Colorado Plateau deposits, where precipitation of metaschoepite from aqueous U concentrations between pH 6 and 10 was reported.²⁴ The equilibrium constants estimated by modeling these batch experiments at circumneutral conditions were then used as a foundation for identifying the reaction rate constants for U–V bearing mineral dissolution in the column experiments.

Column Experiments at pH (7.9)—A linear correlation [with a slope of 1.45 and a R^2 of 0.993] observed between molar concentrations of U and V (Figure S3A) that were mobilized during mine waste reaction with 10 mM HCO₃⁻, suggested dissolution of U–V bearing mineral phases as a potential source. These results agree with a previous study from our research group in which a linear correlation between U and V was observed (slope= 0.636 and $R^2 = 0.95$) in batch experiments reacting mine wastes with 10 mM HCO₃⁻ at pH 8.3.⁶ The U and V release patterns obtained from these experiments were then simulated using PFLOTRAN to further understand the processes affecting the reactive transport of U and V.

Similar to modeling results from the batch experiments, the reactive transport model for flow-through column experiments on reaction with 10 mM HCO₃⁻ suggested dissolution of U–V bearing minerals as the primary mechanism affecting U and V transport. The effluent concentrations of U and V reported as a function of pore volumes (Figure 1C,D) and time (Figure S4), were compared to those obtained from the reactive transport model (PFLOTRAN). PFLOTRAN used $K_{eq} = 10^{-44.81}$ for U–V bearing mineral dissolution estimated from batch experiments by assuming the stoichiometry of carnotite, to adequately fit the flow-through column experiments data and estimate the k_m and fitting parameter (n_m) for U–V bearing mineral phase dissolution. The fitting parameter n_{m} to which the nonlinear water–rock reactions are sensitive (SI eq 8), and k_m were estimated to be 5.2, and 4.8 × 10^{-13} mol cm⁻² s⁻¹, respectively. In addition, the negative reaction rate (I_m) indicated the dissolution of U–V bearing minerals. The k_m estimated for the dissolution of U–V bearing minerals $(4.8 \times 10^{-13} \text{ mol cm}^{-2} \text{ s}^{-1})$ during column experiments was different from that estimated for the batch experiments that is, $k_m = 2.13 \times 10^{-14} \text{ mol cm}^{-2} \text{ s}^{-1}$ (Table 1). The increase in k_m cannot specifically be attributed to a single process. However, the difference in the effective reaction rate constant ($k_{\text{effective}} = k_m \times a_m^0$) estimated using the average surface area, can be attributed to differences in grain size and reactivity conditions, which include flow and heterogeneity in sample composition (Table 1). Observing the effect of grain size on the reactive transport of U and V is also important^{45, 46} as the sieved samples may contain nanocrystals of U-V and other co-occurring U and V minerals, bound electrostatically to the surface of coarser particles.⁴⁷ After the effect of grain size was introduced into PFLOTRAN, by adding U-V bearing minerals of variable sizes and surface areas, their effective reaction rate constants ($k_{\text{effective}}$) were estimated (Table S4). The change in the estimated $k_{\text{effective}}$ for various U–V bearing mineral phases was negligible (Table S4), leading to the inference that the effect of grain size on the reactive transport of U and V is minimal (Figure S5).

Other secondary mineral phases of U such as oxides, phosphates, silicates, and vanadates soluble at circumneutral pH that have not been considered in this model could also contribute to the release of U and V from mine waste.^{32, 41, 48, 49} For instance, a back scatter electron-scanning electron (BSE-SEM) micrograph which shows the association of V with Fe and K was collected (Figure S6). The co-occurrence of U with other elements (e.g., Se and Sr) was also observed using synchrotron μ -XRF mapping (Figure S7). The contribution of these secondary mineral phases can be expected, especially considering (1) the heterogeneity of mine waste, (2) discrepancies between experimental and model U concentrations, and (3) that the early time-behavior of V was higher in the experimental data than the model (see Figure 1D).⁴³ However, after 3–4 h of reaction, the experimental V concentrations agreed well with the model indicating dissolution of U–V bearing minerals. Following the column experiments with 10 mM HCO₃⁻ the mine waste and background soil were sequentially reacted under acidic conditions to further understand the reactive transport of U and V.

Reactive Transport of U and V at Acidic pH

Batch Experiments at pH (3.8)—The reactive transport model for batch reactions with mine waste under acidic conditions (pH 3.8) suggests dissolution of the U–V bearing

mineral phase as a primary contributor to the reactive transport of U and V. We used PFLOTRAN to fit the time behavior of U and V and obtain the equilibrium ($K_{eq} = 10^{-38.65}$) and reaction rate constant ($k_m = 6.4 \times 10^{-14}$ mol cm⁻² s⁻¹) for U–V bearing mineral dissolution by assuming the stoichiometry of carnotite (Figure 2A,B). As observed in the column experiments conducted at circumneutral pH, the estimated K_{eq} under acidic conditions was different from that of synthetic carnotite ($K_{eq} = 10^{-56.38}$).⁴⁴ Additionally, K_{eq} estimated under acidic conditions ($K_{eq} = 10^{-38.65}$) was also different from that obtained at circumneutral conditions ($K_{eq} = 10^{-44.81}$), suggesting the possible dissolution of two different uranyl vanadates. Furthermore, the difference between the K_{eq} of U–V bearing minerals and carnotite at acidic pH can be attributed to difference in reactivity conditions and crystallinity as indicated previously in the reactive transport model at circumneutral pH. We also observed a negative reaction rate (I_m) for U–V bearing minerals that represents their dissolution at acidic pH. The K_{eq} for U–V bearing mineral phase dissolution identified by modeling the batch reaction under acidic conditions, was then used to estimate the reaction rate constants for U–V bearing mineral dissolution during column experiments.

Column Experiments at pH (3.4)—Consistent with the results obtained for experiments at circumneutral pH, the dissolution of U-V bearing minerals was identified as the key process affecting the reactive transport of U and V during column experiments under acidic conditions. A linear correlation (slope of 0.862 and a R^2 of 0.983) between effluent molar concentrations of U and V was also observed in column experiments under acidic conditions (Figure S3B). However, the rates of dissolution obtained for column experiments at acidic conditions are higher compared to those at circumneutral conditions (Table 1). Given that the p K_a of CH₃COOH is 4.74, >90% of CH₃COOH remains protonated. Therefore, a greater contribution from dissolution by acetic acid (protonated form) over complexation or reduction reactions caused by acetate (nonprotonated form) at pH 3.4 is expected.⁵⁰ These results agree with observations made in our previous study at Blue Gap/Tachee, in which linear correlations with a slope of 1.08 and a R^2 of 0.996 between molar concentrations of U and V were observed.⁶ Additionally, 99.1% U and 92.8% V of the total elemental content present in the mine waste was extracted through the sequential reaction of solids in column experiments with 10 mM HCO₃⁻ and CH₃COOH. Note that the acid extractable content of unreacted mine wastes was 1912.12 mg kg⁻¹ U and 858.01 mg kg⁻¹ V (Table S5). The U and V concentrations released during these continuous sequential reactions showed strong agreement with the modeled U and V concentrations, suggesting dissolution of U-V bearing minerals as a primary contributor to the reactive transport of U and V. Different K_{eq} and k_m values for the dissolution of U-V bearing minerals were obtained for mine waste reaction under circumneutral and acidic conditions during flow-through column experiments (Figure 2C,D and Figure S8). The reactive transport model constructed with a known K_{eq} for metaschoepite $(-10^{5.26})^{42}$ and K_{eq} for U–V bearing mineral phase $(-10^{-38.65})$ estimated from a batch reaction under acidic conditions by assuming the stoichiometry of carnotite, was used to identify the k_m constants for dissolution of U–V bearing minerals. The estimated k_m for the U–V bearing mineral phase, 3.2×10^{-13} mol cm⁻² s⁻¹, was slightly different from those estimated for batch reactions ($k_m = 6.4 \times 10^{-14} \text{ mol cm}^{-2} \text{ s}^{-1}$) (Table 1). As discussed previously for column experiments at circumneutral pH (7.9), this decrease in k_m for U–V bearing mineral dissolution cannot specifically be attributed to a single process.

However, the change in the effective reaction rate constant ($k_{effective}$) can be attributed to differences in grain size and reactivity conditions which include flow and heterogeneity in sample composition (Table 1). The effect of grain size was not particularly tested for the sequential reaction of mine waste with CH₃COOH, as the electrostatically bound micrometer sized U–V bearing minerals would already be lost during reaction with 10 mM HCO₃⁻. In addition, the fitting parameter " n_m ", necessary to model the nonlinear dissolution of U–V bearing mineral phase was estimated to be 4.5 (SI eqs 8). Our PFLOTRAN simulations also estimated the reaction rate (I_m) of U–V bearing mineral phase to be negative suggesting dissolution of the U–V bearing mineral phase. However, the reaction rate for metaschoepite and rutherfordine remained unchanged suggesting minimal contribution from metaschoepite and rutherfordine in the reactive transport of U and V. Although, our experimental and model concentrations agree very well with each other, the early release of lower experimental U and V concentrations can be attributed to (1) sample heterogeneity; (2) restricted access to incorporated U–V bearing mineral phases; and (3) precipitation of an unidentified U and V phase.

The difference in the reactive transport of U and V after reaction under circumneutral and acidic conditions could be attributed to the dissolution of different coexisting U–V bearing mineral phases, possibly amorphous and crystalline, as identified in the unreacted mine waste. The coexistence of amorphous and crystalline U-V bearing minerals was identified through selected area electron diffraction (SAED) and energy dispersive spectroscopy (EDS) analysis performed using a high resolution-transmission electron microscopy (HR-TEM) (Figure 3). For instance, Figure 3A,D represents the back scatter electron image (BSE) of two nanoparticulate grains that were identified as uranyl vanadates co-occurring with other minerals based on the EDS elemental composition spectrum in Figure 3B,E. Diffused rings in the SAED pattern (Figure 3C) suggested the identified U-V bearing mineral phase in Figure 3A to be amorphous. However, unlike Figure 3C, definite patterns in Figure 3F suggested the presence of crystalline U-V bearing mineral that co-occurs with other crystalline mineral phases within a particular grain of the mine waste sample. It is likely that the dissolution of the amorphous U–V bearing mineral phase is dominant at circumneutral conditions (pH 7.9) and the dissolution of crystalline U-V bearing mineral phases is dominant under acidic conditions (pH 3.4). However, further analysis is necessary to better understand the effect of minerology, specific stoichiometry, and crystallinity on the reactive transport of U and V from these mine wastes. The coexistence of crystalline and amorphous U-V bearing minerals in unreacted mine waste samples can be a result of nucleation^{51, 52} or polymorphism,⁵³ which are common in natural samples, due to their heterogeneity in site and reactivity conditions. These results agree with observations made in other studies that have shown that crystallinity can affect the solubility of mineral phases.^{54, 55} However, the amorphous characteristics in the U-V bearing mineral phase observed in Figure 3C could also be caused by rapid amorphization due to short-term exposure to the electron beam.^{56, 57} Therefore, our future work will investigate the crystal chemistry of these U-V bearing minerals.

It is possible that other secondary mineral phases such as uranium-bearing oxides, hydrous ferric oxides, and uranium phosphates that have not been considered in the reactive transport model but are soluble at acidic pH, can also affect the reactive transport of U and V.^{42, 58, 59}

For instance, analyses conducted using synchrotron-based micro-XRF mapping on the mine waste sample indicate that U co-occurs with other elements (Figure S7). Despite the chemical complexity of these samples, our reactive transport model can reasonably represent the release of U and V by considering the dissolution of U–V bearing minerals as the key contributing process. However, we acknowledge that contributions from other mineral phases, heterogeneity, and other aqueous processes were not accounted in the reactive transport for U and V, which represents a limitation of this study. Therefore, future research should focus on trying to understand the role of other reaction processes such as the contribution of other secondary mineral/nano-particulate phases of U and V, and the influence of mineral structure of the U–V bearing minerals on the reactive transport U and V from such mine waste sites.

Environmental Implications

This work integrates reactive transport modeling, electron microscopy, and aqueous chemistry methods to evaluate the mechanisms affecting the transport of U and V from mine wastes from a site in northeastern AZ. Outcomes from this investigation include the determination of solubility and reaction rate constants for the dissolution of U–V bearing mineral phases at circumneutral and acidic pH, that can serve as a foundation to better understand their reactivity at relevant field conditions. This information can be useful to better understand the mobility of U and V in neighboring community water sources to assess risks for human exposure. Additionally, the identification of factors affecting the dissolution of U–V bearing minerals under environmentally relevant conditions evaluated in this study is relevant to inform remediation and resource recovery initiatives in sites where these U–V bearing minerals are abundant. For instance, the results from this study have implications for other abandoned uranium mines in the Colorado Plateau, South Dakota (Black Hills), southwest China, southern Jordan, and the calcreted drainages of arid and semiarid western and southern Australia where carnotite, tyuyamunite and other U–V bearing minerals are commonly found.^{24, 46, 60–65}

Spectroscopy and microscopy analyses suggest that U is associated with different elements, which is indicative of the complex mineralogy of these solids. These results are consistent with those reported in the previous study from our research group.⁶ Despite the chemical and mineralogical complexity of these mine wastes, dissolution of U–V bearing minerals was identified as the key mechanism controlling the reactive transport of U and V under the environmentally relevant conditions selected for this study. In addition, the integrated methodology used for this study is transferable to other mining and milling sites, and abandoned U mine waste sites where a better understanding of the reactive transport of U and its co-occurring elements is necessary.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est. 7b03823.

Additional materials and methods, five tables (S1 to S5), and eight figures (S1 to S8) (PDF)

References

- Dawson SE. Navajo Uranium Workers and the Effects of Occupational Illnesses a Case-Study. Hum. Organ. 1992; 51(4):389–397.
- 2. Hopenhayn C. Arsenic in drinking water: Impact on human health. Elements. 2006; 2(2):103-107.
- Byczkowski JZ, Kulkarni AP. Oxidative stress and pro-oxidant biological effects of vanadium. Adv. Environ. Sci. Technol.-New York. 1998; 31:235–264.
- 4. Hund L, Bedrick EJ, Miller C, Huerta G, Nez T, Ramone S, Shuey C, Cajero M, Lewis J. A Bayesian framework for estimating disease risk due to exposure to uranium mine and mill waste on the Navajo Nation. J. R. Statist. Soc. Ser. A. 2015; 178(4):1069–1091.
- Chenoweth, W. Vanadium mining in the Carrizo Mountains, 1942–1947. San Juan County, New Mexico and Apache County Arizona. Vol. 378. New Mexico Bureau of Mines and Mineral Resources; 1991. p. 33Open-file Report
- Blake JM, Avasarala S, Artyushkova K, Ali A-MS, Brearley AJ, Shuey C, Robinson WP, Nez C, Bill S, Lewis J. Elevated concentrations of U and co-occurring metals in abandoned mine wastes in a northeastern Arizona Native American community. Environ. Sci. Technol. 2015; 49(14):8506– 8514. [PubMed: 26158204]
- Qafoku NP, Zachara JM, Liu C, Gassman PL, Qafoku OS, Smith SC. Kinetic desorption and sorption of U(VI) during reactive transport in a contaminated Hanford sediment. Environ. Sci. Technol. 2005; 39(9):3157–3165. [PubMed: 15926566]
- Dong W, Brooks SC. Determination of the Formation Constants of Ternary Complexes of Uranyl and Carbonate with Alkaline Earth Metals (Mg²⁺, Ca²⁺, Sr²⁺, and Ba²⁺) Using Anion Exchange Method. Environ. Sci. Technol. 2006; 40(15):4689–4695. [PubMed: 16913125]
- 9. Bargar JR, Reitmeyer R, Davis JA. Spectroscopic confirmation of uranium (VI)-carbonato adsorption complexes on hematite. Environ. Sci. Technol. 1999; 33(14):2481–2484.
- Naeem A, Westerhoff P, Mustafa S. Vanadium removal by metal (hydr)oxide adsorbents. Water Res. 2007; 41(7):1596–1602. [PubMed: 17303211]
- Burke IT, Mayes WM, Peacock CL, Brown AP, Jarvis AP, Gruiz K. Speciation of arsenic, chromium, and vanadium in red mud samples from the Ajka spill site, Hungary. Environ. Sci. Technol. 2012; 46(6):3085–3092. [PubMed: 22324637]
- Peacock CL, Sherman DM. Vanadium (V) adsorption onto goethite (*a*-FeOOH) at pH 1.5 to 12: a surface complexation model based on ab initio molecular geometries and EXAFS spectroscopy. Geochim. Cosmochim. Acta. 2004; 68(8):1723–1733.

- Catana G, Rao RR, Weckhuysen BM, Van Der Voort P, Vansant E, Schoonheydt RA. Supported vanadium oxide catalysts: Quantitative spectroscopy, preferential adsorption of V^{4+/5+}, and Al₂O₃ coating of zeolite Y. J. Phys. Chem. B. 1998; 102(41):8005–8012.
- 14. Sylwester E, Hudson E, Allen P. The structure of uranium (VI) sorption complexes on silica, alumina, and montmorillonite. Geochim. Cosmochim. Acta. 2000; 64(14):2431–2438.
- Stewart BD, Cismasu AC, Williams KH, Peyton BM, Nico PS. Reactivity of Uranium and Ferrous Iron with Natural Iron Oxyhydroxides. Environ. Sci. Technol. 2015; 49(17):10357–10365.
 [PubMed: 26226398]
- Liu C, Shang J, Shan H, Zachara JM. Effect of subgrid heterogeneity on scaling geochemical and biogeochemical reactions: A case of U(VI) desorption. Environ. Sci. Technol. 2014; 48(3):1745– 1752. [PubMed: 24377314]
- Kosmulski M. pH-dependent surface charging and points of zero charge: III. Update. J. Colloid Interface Sci. 2006; 298(2):730–741. [PubMed: 16466735]
- Schroth BK, Sposito G. Surface charge properties of kaolinite. MRS Online Proc. Libr. 1996; 432:87.
- Bernier-Latmani R, Veeramani H, Vecchia ED, Junier P, Lezama-Pacheco JS, Suvorova EI, Sharp JO, Wigginton NS, Bargar JR. Non-uraninite products of microbial U (VI) reduction. Environ. Sci. Technol. 2010; 44(24):9456–9462. [PubMed: 21069950]
- Hansley PL, Spirakis CS. Organic matter diagenesis as the key to a unifying theory for the genesis of tabular uranium-vanadium deposits in the Morrison Formation, Colorado Plateau. Econ. Geol. Bull. Soc. Econ. Geol. 1992; 87(2):352–365.
- Tokunaga TK, Kim Y, Wan J. Potential remediation approach for uranium-contaminated groundwaters through potassium uranyl vanadate precipitation. Environ. Sci. Technol. 2009; 43(14):5467–5471. [PubMed: 19708383]
- Spano TL, Dzik EA, Sharifironizi M, Dustin MK, Turner M, Burns PC. Thermodynamic investigation of uranyl vanadate minerals: Implications for structural stability. Am. Mineral. 2017; 102(6):1149–1153.
- 23. Franczyk, KJ. Stratigraphic revision and depositional environments of the Upper Cretaceous Toreva Formation in the northern Black Mesa area. Navajo and Apache Counties: AZ; Bulletin 1685: U.S. Geological Survey; 1988.
- Hostetler P, Garrels R. Transportation and precipitation of uranium and vanadium at low temperatures, with special reference to sandstone-type uranium deposits. Econ. Geol. Bull. Soc. Econ. Geol. 1962; 57(2):137–167.
- 25. Giammar DE, Hering JG. Time scales for sorption-desorption and surface precipitation of uranyl on goethite. Environ. Sci. Technol. 2001; 35(16):3332–3337. [PubMed: 11529573]
- Davis JA, Meece DE, Kohler M, Curtis GP. Approaches to surface complexation modeling of uranium (VI) adsorption on aquifer sediments. Geochim. Cosmochim. Acta. 2004; 68(18):3621– 3641.
- 27. Curtis GP, Davis JA, Naftz DL. Simulation of reactive transport of uranium (VI) in groundwater with variable chemical conditions. Water Resour. Res. 2006; 42(4):W04404.
- Hammond GE, Lichtner PC. Field-scale model for the natural attenuation of uranium at the Hanford 300 Area using high-performance computing. Water Resour. Res. 2010; 46(9):W09527.
- Zachara JM, Chen X, Murray C, Hammond G. River stage influences on uranium transport in a hydrologically dynamic groundwater-surface water transition zone. Water Resour. Res. 2016; 52:1568.
- Singer DM, Zachara JM, Brown GE Jr. Uranium speciation as a function of depth in contaminated Hanford sediments-A micro-XRF, micro-XRD, and micro-and bulk-XAFS study. Environ. Sci. Technol. 2009; 43(3):630–636. [PubMed: 19244994]
- 31. Stubbs JE, Veblen LA, Elbert DC, Zachara JM, Davis JA, Veblen DR. Newly recognized hosts for uranium in the Hanford Site vadose zone. Geochim. Cosmochim. Acta. 2009; 73(6):1563–1576.
- Tokunaga TK, Kim Y, Wan J, Yang L. Aqueous uranium (VI) concentrations controlled by calcium uranyl vanadate precipitates. Environ. Sci. Technol. 2012; 46(14):7471–7477. [PubMed: 22734621]
- 33. Langmuir, D. Aqueous Environmental Geochemistry. Prentice Hall; 1997.

- 34. Liu C, Zachara JM, Qafoku NP, Wang Z. Scale-dependent desorption of uranium from contaminated subsurface sediments. Water Resour. Res. 2008; 44(8):W08413.
- 35. Steefel CI, DePaolo DJ, Lichtner PC. Reactive transport modeling: An essential tool and a new research approach for the Earth sciences. Earth Planet. Sci. Lett. 2005; 240(3):539–558.
- 36. Shabansefidan F, Reddy MA. Investigation of Uranium Solubility and its Transport Path in the Razgah Basin's Groundwater in the Northwest of Iran, Using Saturation Indexes. Intl. J. Comput. Sci. Manage. Studies. 2012; 4:39.
- Puigdomenech, I., Bruno, J. Modelling uranium solubilities in aqueous solutions: Validation of a thermodynamic data base for the EQ3/6 geochemical codes. Swedish Nuclear Fuel and Waste Management Company; 1988.
- Kubatko K-A, Helean KB, Navrotsky A, Burns PC. Thermodynamics of uranyl minerals: Enthalpies of formation of rutherfordine, UO₂CO₃, andersonite, Na₂CaUO₂(CO₃).3 (H₂O)₅, and grimselite, K3NaUO₂(CO₃).3H2O. Am. Mineral. 2005; 90(8–9):1284–1290.
- Bond DL, Davis JA, Zachara JM. Uranium (VI) release from contaminated vadose zone sediments: Estimation of potential contributions from dissolution and desorption. Developments in Earth and Environmental Sciences. 2007; 7:375–416.
- 40. USEPA Addressing Uranium Contamination on the Navajo Nation. U.S. Environmental Protection Agency, Pacific Southwest, Region 9; San Francisco, CA: 2015.
- 41. Lollar, BS. Environmental Geochemistry. Vol. 9. Elsevier; 2005.
- Riba O, Walker C, Ragnarsdottir KV. Kinetic studies of synthetic metaschoepite under acidic conditions in batch and flow experiments. Environ. Sci. Technol. 2005; 39(20):7915–7920. [PubMed: 16295856]
- 43. Yoshida S, Iguchi T, Ishida S, Tarama K. Some physico-chemical properties of vanadium oxide supported on silica or γ-alumina. Bull. Chem. Soc. Jpn. 1972; 45(2):376–380.
- 44. Langmuir D. Uranium solution-mineral equilibria at low temperatures with applications to sedimentary ore deposits. Geochim. Cosmochim. Acta. 1978; 42(6):547–569.
- Hochella MF, Lower SK, Maurice PA, Penn RL, Sahai N, Sparks DL, Twining BS. Nanominerals, mineral nanoparticles, and earth systems. Science. 2008; 319(5870):1631–1635. [PubMed: 18356515]
- 46. Kim C, Chi C, Miller S, Rosales R, Sugihara E, Akau J, Rytuba J, Webb S. (Micro) spectroscopic analyses of particle size dependence on arsenic distribution and speciation in mine wastes. Environ. Sci. Technol. 2013; 47(15):8164–8171. [PubMed: 23889478]
- McCarthy JF, Zachara JM. Subsurface transport of contaminants. Environ. Sci. Technol. 1989; 23(5):496–502.
- Um W, Wang Z, Serne RJ, Williams BD, Brown CF, Dodge CJ, Francis AJ. Uranium phases in contaminated sediments below Hanford's U tank farm. Environ. Sci. Technol. 2009; 43(12):4280– 4286. [PubMed: 19603635]
- Delemos JL, Bostick BC, Quicksall AN, Landis JD, George CC, Slagowski NL, Rock T, Brugge D, Lewis J, Durant JL. Rapid dissolution of soluble uranyl phases in arid, mine-impacted catchments near Church Rock, NM. Environ. Sci. Technol. 2008; 42(11):3951–3957. [PubMed: 18589950]
- 50. Goss K-U. The p K a values of PFOA and other highly fluorinated carboxylic acids. Environ. Sci. Technol. 2007; 42(2):456–458.
- Kim K-W, Hyun J-T, Lee K-Y, Lee E-H, Lee K-W, Song K-C, Moon J-K. Effects of the different conditions of uranyl and hydrogen peroxide solutions on the behavior of the uranium peroxide precipitation. J. Hazard. Mater. 2011; 193:52–58. [PubMed: 21802844]
- Schoonen M, Barnes H. Reactions forming pyrite and marcasite from solution: I. Nucleation of FeS₂ below 100°C. Geochim. Cosmochim. Acta. 1991; 55(6):1495–1504.
- 53. Lippmann, F. Sedimentary Carbonate Minerals. Vol. 6. Springer Science & Business Media; 2012.
- Stefánsson A, Gíslason SR. Chemical weathering of basalts, Southwest Iceland: effect of rock crystallinity and secondary minerals on chemical fluxes to the ocean. Am. J. Sci. 2001; 301(6): 513–556.
- 55. Gorman-Lewis D, Burns PC, Fein JB. Review of uranyl mineral solubility measurements. J. Chem. Thermodyn. 2008; 40(3):335–352.

- 56. Rey A, Utsunomiya S, Giménez J, Casas I, de Pablo J, Ewing RC. Stability of uranium (VI) peroxide hydrates under ionizing radiation. Am. Mineral. 2009; 94(2–3):229–235.
- Utsunomiya S, Ewing RC. Radiation-induced decomposition of U (VI) alteration phases of UO₂. MRS Online Proc. Libr. 2006; 932(73.1)doi: 10.1557/PROC-932-73.1
- Smith SC, Douglas M, Moore DA, Kukkadapu RK, Arey BW. Uranium extraction from laboratorysynthesized, uranium-doped hydrous ferric oxides. Environ. Sci. Technol. 2009; 43(7):2341–2347. [PubMed: 19452884]
- Ilton ES, Zachara JM, Moore DA, Mckinley JP, Eckberg AD, Cahill CL, Felmy AR. Dissolution study of metatorbernite: Thermodynamic properties and the effect of pH and phosphate. Environ. Sci. Technol. 2010; 44(19):7521–7526. [PubMed: 20804183]
- 60. Mann, A. Chemical Ore Genesis Models for the Precipitation of Carnotite in Calcrete. Commonwealth Scientific and Industrial Research Organisation; 1974.
- 61. Hillebrand W. Carnotite and tyuyamunite and their ores in Colorado and Utah. Am. J. Sci. 1924; 45:201–216.
- 62. Crook T, Blake G. On carnotite and an associated mineral complex from South Australia. Mineral. Mag. 1910; 15(71):271–284.
- 63. Sharma RK, Putirka KD, Stone JJ. Stream sediment geochemistry of the upper Cheyenne River watershed within the abandoned uranium mining region of the southern Black Hills, South Dakota, USA. Environ. Earth Sci. 2016; 75(9):1–12.
- 64. Xu J, Zhu S-Y, Luo T-Y, Zhou W, Li Y-L. Uranium mineralization and its radioactive decayinduced carbonization in a black shale-hosted polymetallic sulfide ore layer, southwest China. Econ. Geol. Bull. Soc. Econ. Geol. 2015; 110(6):1643–1652.
- 65. Abed AM, Saffarini GA, Sadaqah RM. Spatial distribution of uranium and vanadium in the upper phosphorite member in Eshidiyya basin, southern Jordan. Arabian J. Geosci. 2014; 7(1):253–271.

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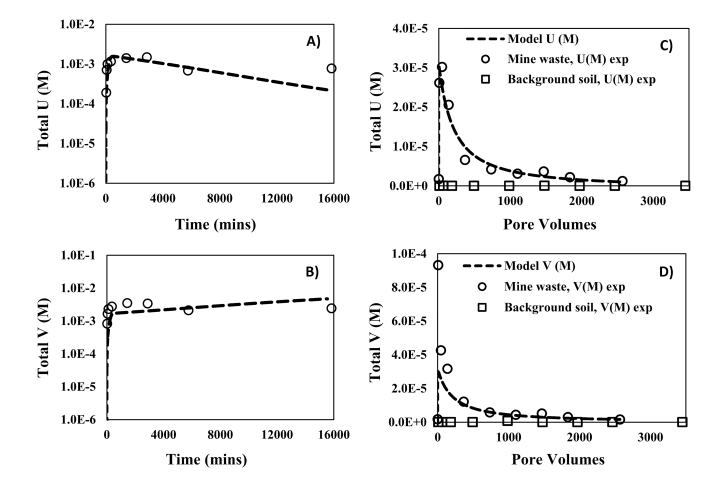


Figure 1.

Measured and simulated effluent concentrations and reactive transport model (PFLOTRAN) of U and V, from mine waste (circle) and background soil (squares) during batch and continuous flow-through column experiments at circumneutral pH (using 10 mM HCO₃⁻). (A) U concentrations from batch experiments versus time; (B) V concentration from batch experiments versus time; (C) U concentrations from column experiments versus pore volumes; (D) V concentration from column experiments versus pore volumes. The curve fitting resulting from the reactive transport model are presented with dashed lines.

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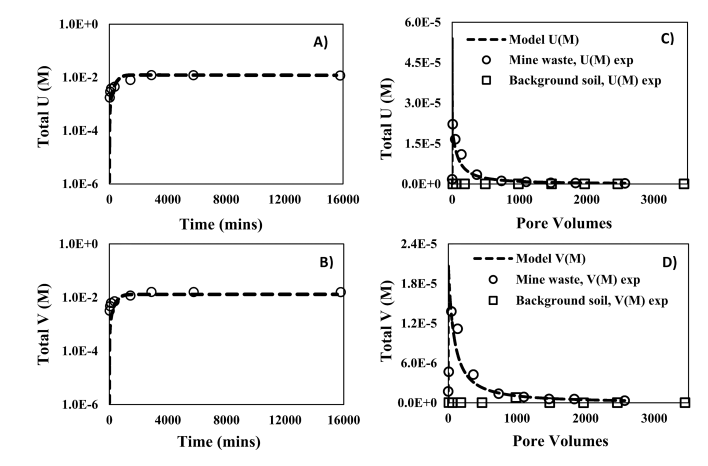


Figure 2.

Measured and simulated effluent concentrations and reactive transport model (PFLOTRAN) of U and V, from mine waste (circle) and background soil (squares) during batch and continuous flow-through column experiments at acidic pH (using 10 mM $C_6H_8O_6$ and CH₃COOH). (A) U concentrations from batch experiments versus time; (B) V concentration from batch experiments versus time; (C) U concentrations from column experiments versus pore volumes; (D) V concentration from column experiments versus pore volumes. The curve fittings resulting from the reactive transport model are presented with dashed lines.

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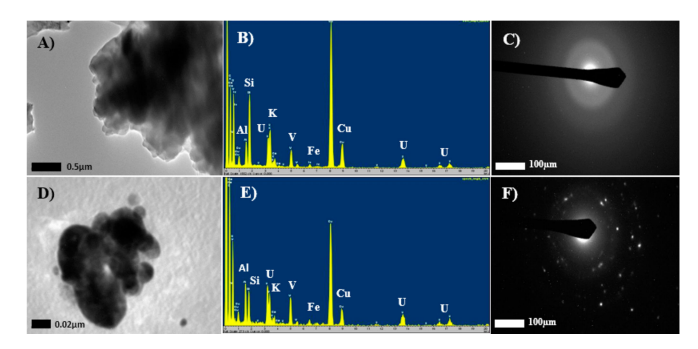


Figure 3.

Transmission electron microscopy (TEM) images, energy dispersive spectroscopy (EDS) spectra and selected area electron diffraction (SAED) patterns for unreacted mine waste samples indicating the co-occurrence of amorphous and crystalline U–V bearing minerals: (A, D) TEM images of U–V bearing mineral phases; (B, E) EDS spectra identifying the presence of U–V bearing minerals; (C, F) SAED patterns confirming the co-occurrence of amorphous and crystalline U–V bearing mineral phase.

Table 1

Parameters of U–V Bearing Minerals Estimated by Modelling the Reactive Transport of U and V during Mine Waste Reaction at Circumneutral and Acidic pH^a

experiments	equilibrium constant (K _{eq})	average U–V bearing mineral surface area $\binom{a_m^0}{(\mathrm{cm}^2\mathrm{cm}^3)}$	reaction rate constants (k_m) (mol cm ⁻² s ⁻¹)	effective reaction rate constant $(k_{\text{effective}} = (k_m \times a_m^0))$ (mol cm ⁻³ s ⁻¹)
batch circumneutral (pH 8.3) (<63 μ m)	10-44.81	468.8	2.13×10^{-14}	9.99×10^{-12}
batch acidic (pH 3.8)(<63 µm)	$10^{-38.65}$	468.8	$6.4 imes 10^{-14}$	$3 imes 10^{-11}$
column circumneutral (pH 7.9) (120– 355 µm)	10-44.81	62.5	4.8×10^{-13}	3×10^{-11}
column acidic (pH 3.4) (120–355 µm)	10 ^{-38.65}	62.5	$3.2 imes 10^{-13}$	2×10^{-11}

^{*a*}The surface area of U–V bearing minerals was estimated using the average U–V bearing mineral diameter (Table S2) and equation 13. The effective reaction rate constant $(k_{\text{effective}} = k_m * a_m^0)$ accounts for the effect of grain surface area on the reactive transport of U and V, where *k* is the reaction rate constant and (a_m^0) is the surface area of U–V bearing minerals.