- 1 Removal of TcO<sub>4</sub> from Representative Nuclear Waste Streams with Layered Potassium Metal *Sulfide Materials*
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# **Abstract:**

Many efforts have focused on the sequestration and immobilization of  $\frac{99}{Tc}$  because the radionuclide is highly mobile in oxidizing environments and presents serious health risks due to 14 its radiotoxicity and long half-life  $(t_{1/2} = 213\,000\,a)$ . One of the more common methods for Tc removal from solution and immobilization in solids is based on reducing Tc from highly soluble 16 Tc<sup>(VII)</sup> to sparingly soluble  $Tc^{(IV)}$ . In order to remove solution Tc through this reduction process, the Tc-sequestering solid must contain a reducing agent and, ideally, the Tc-sequestering material would function in a large range of chemical environments. For long-term stability, the reduced Tc would preferentially be incorporated into the resulting mineral structure instead of simply being sorbed onto the mineral surface. Here, we report results obtained from batch sorption experiments performed in anoxic and oxic conditions with two sulfide-containing

 potassium metal sulfide (KMS) materials, known as KMS-2 and KMS-2-SS. In deionized water 2 in anoxic conditions after 15 d of contact, KMS-2 is capable of removing ~45% of Tc and KMS- 2-SS is capable of removing ~90% of Tc. The improved performance of KMS-2-SS compared to KMS-2 in deionized water in anoxic conditions appears to be linked both to a higher pH resulting from the batch sorption experiments performed with KMS-2-SS and a higher overall 6 purity of KMS-2-SS. Both materials perform even better in highly caustic ( $pH~13.5$ ), high ionic strength (8.0 M) simulated Hanford low-activity waste solutions, removing more than 90% Tc after 15 d of contact in anoxic conditions. Post-reaction solids analysis indicate that Tc(VII) is reduced to Tc(IV) and that Tc(IV) is bonded to S atoms in the resulting KMS-2 structure in a Tc<sub>2</sub>S<sub>7</sub> form. In contrast to previous ion exchange experiments with other KMS materials, the batch sorption experiments examining Tc removal cause the initially crystalline KMS materials to lose much of their initial long-range order.

## 1 **Introduction**

2 The <sup>99</sup>Tc isotope is generated primarily during the fission of <sup>235</sup>U and <sup>239</sup>Pu. Due to its high 3 mobility in oxidizing environments, radiotoxicity, and long half-life  $(t_{1/2} = 213\ 000\ a)$ , 4 management of <sup>99</sup>Tc in radioactive waste streams is of particular importance at nuclear reactors, 5 at nuclear fuel reprocessing facilities such as Sellafield, United Kingdom and La Hague, France, 6 and at legacy waste sites from Cold War activities situated around the world.<sup>1-2</sup> Additionally, the 7 inventory of  $99$ Tc is expected to grow if nuclear power generation is pursued by countries hoping 8 to increase their energy production without increasing their emission of greenhouse gases. 9 Therefore, <sup>99</sup>Tc management is important both in assisting in cleanup efforts of legacy waste 10 sites around the world and ensuring safe disposal of radioactive waste from power generation 11 from nuclear reactors.

12 At one existing legacy waste site, Hanford Site, in southeast Washington State, USA, 13 approximately 1.99 Mg of  $\rm{^{99}Tc}$  (or 1.25 PBq) was produced between 1943 and 1987.<sup>3</sup> The 14 majority of the <sup>99</sup> Tc at the site is located in single- and double-shell tanks containing mixed and 15 radioactive wastes and exists primarily as pertechnetate  $[TeO<sub>4</sub>]$  (from this point forward, we will 16 refer to  $^{99}$ Tc simply as Tc unless specifying the isotope is necessary in the discussion). Current 17 cleanup plans call for separating the tank waste into high-volume, low-activity waste (LAW) and 18 low-volume, high-level waste (HLW) fractions, which will be vitrified at separate facilities in a 19 borosilicate glass matrix. Before vitrification, the aqueous LAW fraction consists of a highly 20 caustic (~pH 13.5), high ionic strength aqueous solution containing  $\mathrm{Na}^+(5-10 \mathrm{M})$ , K<sup>+</sup>, Al(OH)<sub>4</sub><sup>-</sup>, 21 Cl, F, NO<sub>2</sub>, NO<sub>3</sub>, OH, CO<sub>3</sub><sup>2</sup>, organics, and minor ionic species including dissolved metals 22 such as Cr, Ni, Cd, and Pb, and radionuclides. The majority of the  $^{99}$ Tc is expected to partition to 23 the LAW fraction.<sup>4</sup> However, immobilization of Tc in the vitrified product is difficult because Tc

1 volatilizes at the high processing temperatures  $(\sim 1150 \degree C)$  and has relatively low solubility in the 2 borosilicate glass matrix,  $\sim$ 2000 ppm to  $\sim$ 2800 ppm, depending on redox conditions.<sup>5-6</sup> It would, therefore, be advantageous to find a material capable of removing the  $TcO<sub>4</sub>$  from the chemically 4 complex aqueous waste streams before vitrification and capable of immobilizing the Tc in a 5 durable low-temperature waste form for long-term disposal. However, separating the  $TcO<sub>4</sub>$  from 6 the complex aqueous LAW waste stream is particularly challenging due to its high pH and ionic 7 strength.

8 Two techniques that can remove  $TcO<sub>4</sub>$  from aqueous solutions are solvent extraction and 9 sorption (chemisorption and/or physisorption). Solvent extraction, such as the PUREX or UREX 10 processes, is based on an organic phase that strips the  $TcO<sub>4</sub>$  from the aqueous phase.<sup>7-8</sup> In the 11 case of sorption on a material surface, Tc removal is either through an ion exchange process<sup>9-12</sup> 12 or a reductive process whereby the sorbent reduces the highly soluble  $Te^{(VII)}O_4$  to a sparingly 13 soluble Tc(IV) form such as  $Tc^{(IV)}O_2 \cdot nH_2O (2.6 \times 10^{-9} M);^{13}$  the associated redox reaction and 14 standard reduction potential is shown in Equation  $(1)$ .<sup>14</sup>

15 
$$
TCO_4 + 4H^+ + 3e \leftrightarrow TcO_2 + 2H_2O
$$
  $(E = 0.746 \pm 0.012 \text{ V})$  (1)

 A recent review of materials capable of removing Tc from solution identified several 17 materials that reductively separate  $Tc^{15}$ . These materials include apatites,  $12,16$  metals such as  $\frac{\text{zinc}^{17} \text{ or iron}}{\text{.}^{18-20}}$  iron-sulfides,<sup>21</sup> blast furnace slags,<sup>22</sup> chalcogels,<sup>23</sup> and activated carbon,<sup>12,24-27</sup> The effectiveness of these materials is generally determined using batch sorption experiments in 20 anoxic environments. Other than a few studies,  $2^{1,27-28}$  the batch sorption tests are performed in simulated groundwater solution with circumneutral pH and ionic strengths several orders of magnitude lower than that of aqueous LAW. However, to understand the effectiveness of

 materials to remove Tc from aqueous LAW, experiments must be performed at higher pH and ionic strength in the presence of competing ions.



 (XAS) were used to determine the Tc oxidation state using X-ray absorption near edge structure (XANES) spectroscopy and the local environment of Tc was determined by X-ray absorption 3 fine structure (EXAFS) spectroscopy.. These analyses confirm the role of S (as  $S^2$ ) in Tc removal. X-ray diffraction (XRD) of the pre- and post-reaction materials demonstrated that the s crystalline structure was modified during  $TcO<sub>4</sub>$  removal, which is in contrast to the simple ion-exchange mechanism observed with cation removal by KMS materials.

#### 7 **Experimental Section:**

**Synthesis of K<sub>2</sub><sup>***x***</sup>Mg<sub>***x***</sub>Sn<sub>3-***x***</sub>S<sub>6</sub> (** $x = 0.5$ **-1) (KMS-2). Hydrothermal synthesis: Elemental Sn (76)** 9 mmol, 9.0 g), Mg (38 mmol, 0.92 g), S (266 mmol, 8.5 g), K<sub>2</sub>CO<sub>3</sub> (57 mmol, 7.9 g), were mixed in a 23-ml Teflon-lined stainless-steel autoclave. Deionized water (3 mL) was added drop wise until the mixture acquired dough-like consistency. The autoclave was sealed properly and soaked in a preheated oven at 220°C for 15 h under autogenous pressure. Then, the autoclave was allowed to cool at room temperature. A yellow polycrystalline product was isolated by filtration (14.20 g, yield ~70% based on Sn), washed several times with water, acetone, and ether (in that order), and dried under vacuum. Energy dispersive spectroscopy (EDS) analysis gave the 16 average formula "  $K_{1.35}M_{90.6}Sn_{2.6}S_6"$ .

17 **Synthesis of K<sub>2</sub><sup>***x***</sup>Mg<sub>***x***</sub>Sn<sub>3-***x***</sub>S<sub>6</sub> (** $x = 0.5$ **-1) (KMS-2-SS). Solid-state synthesis: A mixture of Sn** 18 (8.9 mmol, 1055 mg), Mg (4.7 mmol, 113 mg), K2S (4.6 mmol, 204 mg), and S (15.7 mmol, 512 19 mg) was sealed under vacuum ( $10^{-4}$  Torr) in a fused silica tube and heated ( $10^{\circ}$ C/h) to 550°C for 20 48 h, followed by cooling to room temperature at  $100^{\circ}$ C/h. The yellow polycrystalline product 21 obtained was washed several times with water, acetone, and ether (in that order)  $(2 g, \sim 85\%)$ 

- 1 yield based on Sn). Energy dispersive spectroscopy (EDS) analysis gave the average formula "K1.3Mg0.95Sn2.1S6".<sup>30</sup> 2
- **Batch Sorption Experiments with KMS-2:** The effectiveness of TcO<sub>4</sub> removal by KMS-2 and
- KMS-2-SS was investigated in 18.2 MQ-cm deionized Millipore<sup>®</sup> water (DIW) and a LAW
- 5 simulant. The LAW simulant is based on the Hanford Tank Waste Operations Simulator
- 6 (HTWOS) model that supports the River Protection Project System Plan Revision  $6.^{38}$  The
- 7 average measured concentration of major elements of the LAW simulant is presented in Table 1.
- 8 The average measured pH of the LAW simulant was 13.5.

9 Table 1 –Measured concentration of major elements of LAW simulant used in tests. The average measured pH of 10 the LAW solution was 13.5.

	<b>Measured</b> concentration			
Constituent	(M)			
Na	8.7			
$\mathbf{A}1$	0.55			
Cr	$3.5 \times 10^{-2}$			
Pb	$4.3 \times 10^{-4}$			
NO <sub>3</sub>	2.5			
NO <sub>2</sub>	0.29			
SO <sub>4</sub>	0.12			
Сl	0.12			

12 Batch sorption tests at  $\sim$ 22 °C involved placing 1.0 g of KMS-2 and KMS-2-SS in contact 13 with 100 mL of solution in a 250-mL polytetrafluoroethylene (PTFE) bottle at room temperature 14 for various durations with periodic solution sampling. Following addition of the DIW or LAW 15 simulant to the bottle, the solutions were spiked with a concentrated stock solution of  $NaTcO<sub>4</sub>$ 16 (0.10 M) to achieve Tc concentrations of  $\sim 5.7 \times 10^{-4}$  M, which is 10× the level predicted by

 HTWOS. Periodic sampling was performed by pipetting a 2-mL aliquot of solution from the test vessels which was immediately filtered (0.2 µm). The volume removed during sampling was not replaced. Experiments were performed either in a fume hood in open atmosphere (oxic) or in a 4 chamber containing N<sub>2</sub> with H<sub>2</sub> (0.7%) to maintain anoxic conditions. The aliquot samples were 5 acidified with 20  $\mu$ L of 70% HNO<sub>3</sub> for analysis with inductively coupled plasma mass spectroscopy (ICP-MS) for Tc and ICP optical emission spectroscopy (ICP-OES) for Cr.

 **Reduction Capacity**: The Ce(IV) method was used to calculate reductive capacity of the KMS-2 8 and KMS-2-SS. The method has previous been reported,  $39-41$  and was applied here with slight modifications. Here, 0.01 g aliquots of the KMS materials were placed in 15 mL of a 40 mM 10 Ce(IV) stock solution prepared with  $10\%$  H<sub>2</sub>SO<sub>4</sub>. The mixture was agitated on a mechanical 11 shaker for 72 h. Following agitation, a 5-mL aliquot was filtered with a 0.45-um syringe filter 12 and titrated with a 20 mM ( $NH_4$ )<sub>2</sub>Fe( $SO_4$ )<sub>2</sub> solution using 0.05 mL of ferroine solution (Fluka) as an indicator. The titration was carried out until a faint pink/blue (lilac) color was observed. The reductive capacity was calculated via the difference between the oxidizing equivalents in the 15 initial Ce(IV) solution and the reducing equivalent of  $Fe(II)$  titrated to react with the excess Ce(IV) remaining following reaction with the KMS-2; the reductive capacity was then corrected for the mass of the sample added to the Ce(IV) solution.

 **X-ray Diffraction:** Powders were loaded into a zero-background holder and diffraction data collected with a Rigaku Miniflex II Bragg-Brentano diffractometer using Cu-Kα radiation  $(\lambda = 1.5418 \text{ Å})$  and a graphite post-diffraction monochromator. Phases were identified by comparing experimental patterns with those simulated from crystal structures using the program TOPAS (v4.2, Bruker AXS). The structure for KMS-2 was obtained from Mertz et. al (2013);

1 other structures were obtained from the ICSD database (Fachinformationszentrum, Karlsruhe, 2 Germany).

 **X-ray Absorption Spectroscopy:** X-ray absorption spectroscopy (XAS) data were obtained at the Stanford Synchrotron Radiation Lightsource Beamline 11-2. The monochromator was detuned 50% to reduce the harmonic content of the beam. Transmission data was obtained using Ar filled ion chambers. Fluorescence data were obtained using a 100 element Ge detector and have been corrected for detector dead time. Data were converted from raw data to spectra using 8 SixPack.<sup>42</sup> Spectra were normalized using Athena.<sup>43</sup> Normalized XANES spectra were fit using standard spectra in the locally-written program, *fites*, which performed non-linear least squares 10 fit of the data.<sup>44</sup> XANES standard spectra were carefully energy calibrated using  $TcO<sub>4</sub>$  adsorbed on Reillex-HPQ as the energy reference. The XANES spectral resolution is 7 eV based on the 12 width of the TcO<sub>4</sub> pre-edge peak, so each spectrum possesses 17 independent data points (range of the spectrum/resolution).

14 To match energy resolution of the  $Tc_2S_7$  spectrum,, XANES spectra of the samples and of 15 the TcO<sub>4</sub><sup>-</sup> reference were convolved with a 1.8 eV Gaussian, and the XANES spectrum of 16  $[(EDTA)Tc(\mu-O)]_2 (Tc^{(IV)}EDTA)$  was convolved with a 1 eV Gaussian.  $Tc^{(IV)}EDTA$  is the 17 Tc(IV) reference spectrum for these samples.  $TcO_2 \cdot xH_2O$  was initially used; however, the 18 spectrum of  $Tc^{(IV)}$ EDTA better matches the sample XANES spectra. The main difference 19 between  $TcO_2 \cdot xH_2O$  and  $Tc^{(IV)}EDTA$  is that the local environment of  $Tc^{(IV)}EDTA$  is more 20 distorted, which makes it a better reference standard for Tc(IV) adsorbed on a mineral surface. 21 Note that  $Tc_2S_7$  is also a  $Tc(IV)$  compound, but its XANES spectrum is distinct from either  $T_{\rm c}$  TcO<sub>2</sub>•*x*H<sub>2</sub>O or Tc<sup>(IV)</sup>EDTA. Tc<sub>2</sub>S<sub>7</sub> may be better formulated as Tc<sub>2</sub>S(S<sub>2</sub>)<sub>3</sub> where S<sub>2</sub> is the 23 disulfide anion,  $S_2^2$ , and the structure is given elsewhere.<sup>45</sup> EXAFS data were fit using Artemis

1 with theoretical scattering curves calculated with Feff 8.5 from the structures of  $Tc_2S_7$  and 2 TcO<sub>2</sub>•*x*H<sub>2</sub>O previously reported.

## 3 **Results and Discussion:**

 *Tc removal from DIW and LAW simulant:* Results of anoxic and oxic removal of Tc by KMS-2 and KMS-2-SS in DIW and the LAW simulant batch experiments as a function of contact time are presented in Figure 1. For the DIW experiments in Figure 1a, it is seen that, apart from the experiment performed in oxic conditions with KMS-2, the Tc concentration decreases with time. It also appears that a maximum removal value was achieved after the 14-d experiment. For batch experiments at 14 d in anoxic conditions with DIW, KMS-2 removal was 10 24 µmol Tc per gram (45% of the available Tc) and KMS-2-SS removal was 50 µmol Tc per gram (92% of the available Tc).



13 Figure 1 – The Tc concentration (µmol/L) as a function contact time with KMS-2 and KMS-2-SS in oxic and anoxic 14 conditions in 1) DIW and 2) the LAW simulant. The error bars (given as 1σ) are the standard deviation of the Tc 15 concentration from duplicate samples. The measured final pH presented for DIW samples is placed near the

 experiment to which is corresponds. The pH for the LAW simulant was invariant for each experiment and is presently merely as the measured average.



20 competition between  $TcO<sub>4</sub>$  and  $O<sub>2</sub>$  for reduction.

 Figure 1b presents the results from the batch sorption experiments performed in LAW simulant. It can clearly be seen that the KMS-2 is more effective in the LAW solution compared with the DIW solution while KMS-2-SS performs slightly better in the LAW simulant compared

 with DIW. In anoxic conditions, KMS-2 removes 56 µmol Tc per gram (95% of the available Tc) from the LAW simulant, and KMS-2-SS removes 59 µmol Tc per gram (97% of the available Tc).

 In oxic conditions, KMS-2 removes 52 µmol Tc per gram (88% of the available Tc), from the LAW solution, and KMS-2-SS removes 58 µmol Tc per gram (97% of the available Tc). The values of more than 50 µmol Tc per gram for KMS-2 in the oxic and anoxic experiments are up to two times the removal compared with batch experiments performed in DIW. The two properties of the LAW simulant that are attributed to the improvement in the Tc removal efficiency of KMS-2 and KMS-2-SS are the higher pH (13.5) and high ionic strength (6 – 8 M Na). Increasing the solution alkalinity has been shown in previous work to improve the reduction of Tc(VII) by sulfide in solution. Increasing the ionic strength of solution has also been shown in 12 previous works to improve the rate and extent of  $Tc(VII)$  removal from solution.<sup>28,46</sup> The results from these batch sorption tests are promising due to the Tc removal capacity of the materials in the highly caustic, high-ionic strength LAW. In a previous study where a set of promising Tc sequestering materials were evaluated, every sorbent except for KMS-2 16 experienced a large decrease in Tc removal capacity in LAW simulant compared to DIW.<sup>47-48</sup>

The decrease in performance by these materials in the LAW simulant was attributed to a

combination of interference from the excess Cr in the simulant solution and the high pH. The

19 initial Cr concentration in the LAW was measured to be  $3.5 \times 10^{-2}$  M. However, the KMS-2 can

from LAW simulant in the oxic batch experiments is shown. At 14 d, the KMS-2 removed 1581

overcome the competitive Cr(VI) effect as demonstrated in Figure 2, where the removal of Cr

22 umol Cr  $g^{-1}$  (47% of the available Cr) and KMS-2-SS removed 1385 umol Cr  $g^{-1}$  (41% of the

- 1 available Cr. This change in Cr levels is also observed by the color change of the solution from
- 2 yellow initially due to Cr(VI) to blue at 14 d due to Cr(III).



3

4 Figure 2 – The Cr concentration (mmol/L) as a function contact time with KMS-2 and KMS-2-SS in oxic conditions 5 in the LAW simulant. The error bars (given as 1σ) are the standard deviation of the Cr concentration from duplicate 6 samples. The pH for the LAW simulant was invariant for each experiment and is presently merely as the measured 7 average.

The ability to remove both Tc and Cr is likely a result of the  $S<sup>2</sup>$  content of the KMS-2 where 9 the  $S^2$  can reduce both Tc(VII) and Cr(VI) to Tc(IV) and Cr(III), respectively. In a measurement 10 of the reductive capacity of the KMS-2 materials, following a method similar to Um et al.,<sup>41</sup> 11 KMS-2 had a reductive capacity of 7.4  $\pm$  0.6 meg g<sup>-1</sup> and the KMS-2-SS measured 20  $\pm$  3 meg g<sup>-1</sup> 12 <sup>1</sup>. These values are equivalent to removing 2.5 mmol  $g^{-1}$  KMS-2 or 6.5 mmol  $g^{-1}$  KMS-2-SS of 13 Tc or Cr.

 *Change in KMS structure:* The XRD patterns for KMS-2 are presented in Figure 3 for the unreacted material, the KMS-2 material after Tc removal from DIW in oxic conditions, and the KMS-2 material after Tc removal from the LAW simulant in oxic conditions. The unreacted 4 material contains substantial amounts of magnesite  $(MgCO<sub>3</sub>)$ , schoenfliesite  $[MgSn(OH)<sub>6</sub>]$ , and 5 SnO<sub>2</sub>. The material has a layered structure as indicated by the large peak at 8.7 Å.



#### 

 Figure 3 – X-ray diffraction patterns of starting material KMS-2 (black pattern), the KMS-2 material after Tc 8 removal from DIW in oxic conditions (green pattern), and the KMS-2 material after Tc removal from LAW 9 simulant in oxic conditions (blue pattern). The  $(M)$  indicates peaks associated with magnesite  $(MgCO<sub>3</sub>)$ , the  $(H)$ 10 indicates  $MgSn(OH)_6$ , the (T) indicates  $SnO_2$ , and the (S) indicates elemental Sulfur.

 For the layered material used in the DIW experiment, the pre- and post-reaction patterns are nearly identical apart from the disappearance of the peak at 8.7 Å. This would indicate that the 13 layered material is largely unaltered upon removal of  $TcO<sub>4</sub>$  from solution. For the sample obtained from the LAW simulant experiment, compared to the largely crystalline starting material, the reacted sample appears to have undergone a major structural change with only

 small, broad peaks associated very small crystallites (nanoparticles). This result is in contrast to previous studies on contaminant removal with KMS-2, which demonstrated ion-exchange 3 behavior where cations from solution replace  $K^+$  in the interlayers on the KMS-2 structure.<sup>29</sup> The great change in the XRD pattern after contact with the LAW simulant suggests that the structure of the KMS-2 material is greatly altered during Tc removal. This alteration could be caused by the loss of sulfide ions from the octahedral layers of the KMS-2.

 *Tc oxidation state and environment in KMS:* The Tc K-edge XAS spectra of the reference standards were obtained and the results are given in Figure S1 (SI). Three standard spectra were 9 used: TcO<sub>4</sub>, Tc<sub>2</sub>S<sub>7</sub>, and Tc<sup>(IV)</sup>EDTA complex. The latter represents Tc(IV) coordinated by oxygen atoms in a distorted octahedral environment. The bulk XANES spectra for KMS-2 and KMS-2-SS samples that were used for Tc removal in anoxic DIW systems are presented in Figure 4. Corresponding XANES fitting parameters are presented in Table 2. For Tc-loaded KMS-2 in anoxic DIW, the major signal contribution arises from a Tc(IV) coordinated by 14 oxygen atoms with a small contribution from  $TcO<sub>4</sub>$ , and, no  $Tc<sub>2</sub>S<sub>7</sub>$  was observed . The KMS-2 had the lowest Tc removal from the DIW batch experiments, and this XANES result illustrates the importance of sulfide in the immobilization of Tc by these materials. For the KMS-2-SS from anoxic DIW batch experiments the contribution to the spectrum comes entirely (within error) 18 from Tc<sub>2</sub>S<sub>7</sub>. The Tc<sub>2</sub>S<sub>7</sub> is a commonly reported compound with Tc(IV) coordinated primarily by 19 disulfide ligands.<sup>45</sup> The KMS-2-SS had higher Tc removal in DIW compared with KMS-2, further demonstrating the importance of sulfide for Tc immobilization.

 Table 2 – XANES fitting results for Tc in KMS-2 in various experimental conditions. The numbers in parentheses are the standard deviation of the contribution of that component for the last digit. The value of p is the probability

1 that the improvement to the fit from including this spectrum is due to noise. Components with  $p < 0.05$  are 2 significant at the 2 $\sigma$  level and those with  $p < 0.01$  are significant at the 3 $\sigma$  level.

<b>Material</b>	<b>Solution</b>	<b>Conditions</b>	TeO <sub>4</sub>	p	Tc(IV)	p	$Tc_2S_7$	p
$KMS-2$	<b>DIW</b>	anoxic	0.09(2)	0.123	0.95(4)	< 0.001	0.00(7)	
$KMS-2-$ SS	DIW	anoxic	0.00(1)		0.00(2)		1.00(3)	$\leq 0.001$
$KMS-2-$ SS	<b>DIW</b>	anoxic to OX1C	0.00(2)	1	0.03(3)		0.97(4)	< 0.001
$KMS-2$	LAW simulant	anoxic	0.00(1)		0.01(2)		0.99(3)	$\leq 0.001$
$KMS-2$	LAW simulant	<b>OXIC</b>	0.25(2)	< 0.001	0.12(3)	< 0.001	0.64(3)	< 0.001

 The Tc<sub>2</sub>S<sub>7</sub> that was formed in the anoxic DIW experiment with KMS-2-SS showed high stability upon exposure to an oxic environment. Figure 4 displays the XANES spectrum and resulting fits from KMS-2-SS that were collected from a batch experiment where the container was transferred to an oxic environment after 14 d of contact in anoxic conditions. Following 7 exposure to oxic conditions, the  $Tc_2S_7$  remained the only Tc species observed by XANES.



1 Figure 4 – Technetium XANES spectra of Tc in the following batch experiments: KMS-2 in DIW and anoxic 2 conditions, KMS-2-SS in DIW and anoxic conditions, and KMS-2-SS in DIW and anoxic conditions followed by 3 oxic conditions.

4 In addition to the major differences that may be observed from the XANES spectra, the local 5 environment of Tc in these samples was determined by EXAFS. The Tc K-edge EXAFS data are 6 presented for selected samples in Figure 5. For the KMS-2 sample in anoxic conditions in DIW, 7 the data are consistent with Tc-S and Tc-Tc distances suggest the presence of  $Tc_2S_7$ .<sup>45</sup> In addition, a small amount of  $TcO<sub>4</sub>$  is present in this sample. For the KMS-2-SS samples from the 9 anoxic and anoxic-to-oxic conditions, the EXAFS spectra and fit parameters are nearly identical. 10 The interatomic distances are slightly different from those previously reported for  $Tc_2S_7$ ,  $Tc-S =$ 11 2.378(2) Å and Tc-Tc = 2.774(2) Å.<sup>45</sup> The local environment of Tc is identical to that in Tc<sub>2</sub>S<sub>7</sub> 12 within error. In general, the KMS materials appear to act as a source for sulfide to reduce  $TcO<sub>4</sub>$ 13 and precipitate it as  $Tc_2S_7$ , which is stable in oxic conditions, at least over short time periods.



1 Figure 5 - EXAFS spectrum (red) and fit (black) (left panel) and Fourier Transform (right panel) for (a) KMS-2

- 2 (anoxic, DIW), (b) KMS-2-SS (anoxic, DIW), and (c) KMS-2-SS (anoxic, exposed to air, DIW). Fitting parameters
- 3 are given in Table 3.|



4 Table 3 - EXAFS Fitting parameters for Figure 5.



5 a) The value of p is the probability that the improvement to the fit from including this spectrum is due to noise. Components with  $p < 0.05$  are significant at the 2 $\sigma$  level and those 7 with  $p < 0.01$  are significant at the 3 $\sigma$  level

8 Bulk XANES spectra were also obtained for KMS-2 from batch experiments performed in

9 anoxic and oxic conditions with the LAW simulant are presented in Figure 6 with the results of

10 the XANES fitting presented in Table 2. Both the spectra from the anoxic and oxic KMS-2

11 experiments show fits in excellent agreement with the data from the standards. The KMS-2 from

12 the anoxic LAW simulant experiment consists primarily of a  $Tc^{(IV)}{}_{2}S_{7}$  species (Figure 6a).

13 However, as seen in Table 2 and Figure 6b, for the KMS-2 sample from the LAW simulant batch

14 experiment performed in oxic conditions, 64% of the Tc contribution is  $Tc^{(IV)}{}_{2}S_7$ , 12% of the

15 contribution is from Tc(IV) coordinated by oxygen atoms, and 25% is unreduced  $Te^{(VII)}O_4$ .

16 These data suggest that effective removal of Tc is possible by KMS-2 in oxidizing conditions,

17 although the reduction reaction from the highly soluble Tc(VII) to the sparingly soluble Tc(IV)

18 may not proceed as quickly as when Tc removal is performed in anoxic conditions due to either

1 competition between  $TcO_4$  and  $O_2$  for reduction by sulfide or due to oxidation of the sulfide 2 ligands of  $Tc_2S_7$  to yield  $Tc(IV)$  species. The EXAFS analyses were not performed on the LAW simulant samples. Data were not obtained from KMS-2-SS samples used for Tc removal from the LAW simulant.



 Figure 6 – Technetium XANES spectra for fits of Tc in the following batch experiments of a) KMS-2 in LAW and anoxic conditions, and b) KMS-2 in LAW and oxic conditions.

#### **Conclusions:**

 The KMS-2 and KMS-2-SS were successful in removing Tc(VII) from a high ion strength, high pH LAW simulant. The Tc(VII) removal from LAW was improved compared with Tc removal from DIW. The ability of the sulfide-containing material to remove Tc increases with increasing pH and ionic strength. This property is unique among Tc sequestering materials where performance is general worse in the highly caustic, high ionic strength LAW simulant. The XRD 14 analyses of the reacted material show that contact with Tc containing LAW greatly alters its structure. XRD patterns of samples that were used to remove Tc from the LAW displayed broad,



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2 **Figure S2. KMS-2 (anoxic, DIW) EXAFS spectrum (red) and fit (black) (left panel) and its Fourier**  3 **Transform (right panel). Fitting parameters are given in Table S1.**

4

# 5 **Table S1. EXAFS Fitting parameters for Figure S2.**



6 a)  $S_0^2 = 0.8$  (fixed),  $\Delta E = 3(2)$  eV;

2 b)  $S_0^2 = 0.8$  (fixed),  $\Delta E = 3(2)$  eV;<br>
3 b) Lukens, W. W.; Bucher, J. J.; Shuh, D. K. Edelstein, N. M.; *Env. Sci. Tech.* **2005**, 39, 8064.<br>
2 b) Parameter fixed at this value

- c) Parameter fixed at this value
- 9



3 **Figure S3. KMS-2-SS (anoxic, DIW) EXAFS spectrum (red) and fit (black) (left panel) and its**  4 **Fourier Transform (right panel). Fitting parameters are given in Table S2.**



Tc  $1.6(5)$   $0.004(1)$   $2.785(6)$   $\leq 0.001$ 



a) Fit range: 2<k<14, 1<R <3; 16.9 independent data. 7 parameters,  $S_0^2 = 0.8 \Delta E_0 = 5(1)$ 8  $eV, \chi^2 = 106, \chi^2 = 13, R = 0.013.$ 

9 b) The value of p is the probability that the improvement to the fit from including this 10 spectrum is due to noise. Components with p < 0.05 are significant at the 2σ level 11 and those with  $p < 0.01$  are significant at the 3 $\sigma$  level.

12

2

5

1





2 **Figure S4. KMS-2-SS (anoxic, exposed to air, DIW) EXAFS spectrum (red) and fit (black) (left**  3 **panel) and its Fourier Transform (right panel). Fitting parameters are given in Table S3.**





6 **a**) Fit range:  $2 \le k \le 14$ ,  $1 \le R \le 3$ ; 16.9 independent data. 7 parameters,  $S_0^2 = 0.8$ ,  $\Delta E_0 = 4(1)$ 7  $eV, \chi^2 = 187, \chi^2 = 23.6, R = 0.020.$ 

8 b) The value of p is the probability that the improvement to the fit from including this 9 spectrum is due to noise. Components with p < 0.05 are significant at the 2σ level 10 and those with p < 0.01 are significant at the 3σ level.