#### **Examination of Uranium(VI) Leaching During Ligand Promoted Dissolution** 1 2 of Waste Tank Sludge Surrogates

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- 4 Brian A. Powell<sup>\*a,\$</sup>, Linfeng Rao<sup>a</sup>, Kenneth. L. Nash<sup>b</sup>
- <sup>a</sup>Lawrence Berkeley National Laboratory, Berkeley, CA 94720 USA
- 5 6 <sup>b</sup>Washington State University, Pullman, WA 99164 USA
- 7 \*Corresponding author: email: bpowell@clemson.edu
- 8 <sup>\$</sup>Current Address: Department of Environmental Engineering and Earth Sciences, Clemson University,
- 9 Anderson, SC 29630
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11	<b>Summary.</b> The dissolution of synthetic boehmite ( $\gamma$ -AlOOH) by 1-hydroxyethane-1,1-
12	diphosphonic acid (HEDPA) was examined in a series of batch adsorption/dissolution experiments.
13	Additionally, the leaching behavior of <sup>233</sup> U(VI) from boehmite was examined as a function of pH and
14	HEDPA concentration. The results are discussed in terms of sludge washing procedures that may be
15	utilized during underground tank waste remediation. In the pH range 4 to 10, complexation of Al(III) by
16	HEDPA significantly enhanced dissolution of boehmite. This phenomenon was especially pronounced in
17	the neutral pH region where the solubility of aluminum, in the absence of complexants, is limited by the
18	formation of sparsely soluble aluminum hydroxides. At pH higher than 10, dissolution of synthetic
19	boehmite was inhibited by HEDPA, likely due to sorption of Al(III):HEDPA complexes. Addition of
20	HEDPA to equilibrated U(VI)-synthetic boehmite suspensions yielded an increase in the aqueous phase
21	uranium concentration. Partitioning of uranium between the solid and aqueous phase is described in terms
22	of U(VI):HEDPA speciation and dissolution of the boehmite solid phase.

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#### 24 **1. Introduction**

25 The underground storage tanks at the Hanford Site in Washington State, U.S.A. contain the byproducts

26 from a number of spent nuclear fuel reprocessing processes including the bismuth phosphate ( $BiPO_4$ ),

27 Redox and PUREX processes (1). Over time the waste has stratified into a salt cake, a supernatant phase,

28 and an underlying sludge phase. Insoluble aluminum oxides make up a significant fraction of the sludge

- 29 phase (1). Most of the transuranics have partitioned into the sludge phase, making vitrification of the
- 30 sludge phase for geologic disposal a plausible treatment process (1). However, vitrification will be

prohibitively expensive due to the large volume of the sludge phase. Therefore, reduction of the volumeof the sludge through dissolution of the aluminum oxides presents a favorable alternative.

33 Various strategies of sludge leaching have been proposed and tested with sludge stimulants, 34 including leaching with increasingly aggressive procedures (0.01M NaNO<sub>2</sub> + 0.01M NaOH, 3M NaOH, 35 0.05M glycolic acid + 0.10M NaOH, 0.10M HNO<sub>3</sub>, 2.0M HNO<sub>3</sub>, 0.5M HEDPA (1-hydroxyethane-1,1-36 diphosphonic acid)) (1-3). It was found that the aluminosilicates cannot be removed using the baseline 37 washing procedure (0.01 M NaNO<sub>2</sub> + 0.01 M NaOH) and no single treatment achieved complete 38 dissolution (1-3). Data from these experiments suggest that HEDPA could be an effective leachant to 39 reduce the volume of waste sludge. In order to develop an efficient waste treatment process, further 40 studies are needed to understand the extent of sludge phase dissolution and the partitioning of actinides 41 (U, Np and Pu) during the leaching process.

42 As a diphosphonic acid, HEDPA is known to form strong complexes with metal ions including 43 actinides and aluminum across a wide pH range (4-6). In the absence of complexing ligands, aluminum 44 (hydro)oxides are only sparingly soluble. Therefore, formation of Al(III):HEDPA complexes will 45 promote sludge dissolution. However, partitioning of the actinides between the sludge and aqueous phase 46 must also be understood in order to evaluate the viability of HEDPA for sludge washing. Formation of 47 U(VI):HEDPA complexes affects the partitioning of uranium during the leaching of waste sludge with 48 HEDPA. Previous experiments demonstrated that greater than 95% of the total uranium was leached from 49 BiPO<sub>4</sub>, Redox, and PUREX sludge waste simulants by washing the sludge with 0.5 M HEDPA (3). 50 Recently, a number of  $(UO_2)_m H_h L_l$  (where L stands for the fully deprotonated HEDPA anion) complexes, 51 ranging from cationic to anionic, were identified in the pH region from 2 to 12 using potentiometry, calorimetry, and spectroscopic techniques (4). The thermodynamic data provided by Reed et al. (4) are 52 53 used below to describe observed behavior of uranium in synthetic boehmite suspensions amended with 54 HEDPA.

55 The objective of this study was to investigate the ability of HEDPA to accomplish dissolution of 56 synthetic boehmite and to examine the leaching of uranium during dissolution. A companion study was also performed to examine the leaching of neptunium and plutonium under similar conditions (7). The
data are expected to assist in the development of remediation strategies to be used during waste tank
sludge washing.

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# 61 **2. Materials and Methods**

# 62 2.1 Solid Phase Characterization

63 The alumina used in this work was obtained from SASOL (trade name CATAPAL® B). Powder X-ray 64 diffraction data, determined using a Seimens D-500 Diffractometer, indicate that the material has the 65 crystal structure of boehmite ( $\gamma$ -AlOOH) although a significant amorphous character was indicated through broad peaks in the XRD spectra. A surface area of 354 m<sup>2</sup> g<sup>-1</sup> was measured by  $N_2(g)$  adsorption 66 67 using a Micrometrics BET Surface Area Analyzer. Potentiometric tirations were conducted with  $100 \text{ g L}^{-1}$ 68 boehmite suspensions in 0.01, 0.1, and 1.0 M NaCl to determine the point-of-zero-salt-effect (pzse). The 69 titration results are shown in Figure 1. The boehmite surface has a net positive or negative charge due to 70 protonation or deprotonation of surface hydroxyl groups with changing pH. The intersection of the 71 curves for the three NaCl concentrations at pH 8.1 represents the point at which changes in the 72 concentration of the background electrolyte have no effect on the net surface charge density (8). This 73 value is consistent with the zero-point-of-charge measured for several synthetic aluminas (9-11). The 74 apparent proton surface charge density was calculated assuming a surface site density of 1.7 sites per nm<sup>2</sup>. 75

# 76 2.2 Preparation of Uranium Working Solution

A stock solution of <sup>233</sup>U(VI) tracer was obtained from the inventory at Lawrence Berkeley National
Laboratory and purified by ion exchange. Analysis by α-spectroscopy indicated that the purified uranium
contains 96.6% <sup>233</sup>U and 3.3% <sup>232</sup>U. A 1.1 x 10<sup>-4</sup> M U(VI) working solution in 1M NaCl at pH 3 was
prepared from the stock solution and used for sorption experiments. The concentration of uranium was

determined by liquid scintillation counting (LSC) using EcoLumeTM (MP Biomedicals Inc.) cocktail on
a Wallac 1415 liquid scintillation counter.

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# 84 2.3 Dissolution of boehmite using HEDPA

85 Working suspensions spanning the pH range 4 to 11 were prepared by equilibrating boehmite with 1.0 M 86 NaCl then using these suspensions to prepare samples for batch experiments. The batch experiments were 87 performed in polypropylene centrifuge tubes shaken along their longitudinal axis on an orbital shaker at 88 150 rpm. Batch dissolution experiments to examine the dissolution of boehmite by HEDPA were conducted in suspensions with a constant boehmite concentration of 600 mg L<sup>-1</sup> (10 mM as Al) containing 89 90 5 mM, 10 mM, and 50 mM HEDPA. At specified intervals, aliquots were removed and passed through 91 200 nm nylon filters (Gelman Acrodisc). The aluminum concentration in the filtrate was measured using 92 ICP-OES (Perkin Elmer, Optima 5300 DV). The pH of each suspension was adjusted towards an initial 93 pH during sampling intervals using 0.1 and 0.01 M HCl and NaOH. At a few intervals, the HEDPA 94 concentration in the filtrate was measured using a standard spectrophotometric method (12) employing a 95 Cary 5G Spectrophotometer.

96 A series of potentiometric titrations were attempted to examine Al(III):HEDPA complex 97 formation in 1 M NaCl. However the titrations were unsuccessful as a white precipitate was observed 98 across the pH range 4 to 9. To identify the precipitate, a series of solutions were prepared in 1 M NaCl 99 with total Al(III):HEDPA ratios varying from 1:1 to 1:5. The precipitates were isolated via centrifugation 100 and re-dissolved in ultra pure H<sub>2</sub>O. The aluminum and HEDPA concentrations in the resulting solution 101 were used to calculate the Al(III):HEDPA molar ratio in the solids. Additionally, to examine the effect of 102 the background electrolyte and electrolyte concentrations, Al(III):HEPDA precipitates were formed in 0.1 103 and 1.0 M NaCl, KCl, NaNO<sub>3</sub>, and (CH<sub>3</sub>)<sub>4</sub>NCl as well as ultra pure water. Aliquots of each solution were 104 passed through filters with nominal pore sizes ranging from 450 nm to 12 nm to determine the particle 105 size range of the precipitates.

#### 107 **2.4 Sorption of uranium to boehmite**

108 The effect of HEDPA on uranium partitioning to synthetic boehmite was examined by amending 109 equilibrated U(VI)-boehmite suspensions with HEDPA as described below. The uranium-boehmite 110 suspensions were prepared by adding an aliquot of the U(VI) working solution to a boehmite suspension previously adjusted to a desired pH. The initial solution conditions were 6.1 µM U(VI), 660 mg L<sup>-1</sup> 111 112 boehmite, and 1.0 M NaCl. The samples were mixed at 150 rpm for 10 days. Preliminary kinetic tests (in 113 the absence of HEDPA) indicated steady state uranium partitioning was achieved after 7 days. At 114 specified intervals, the soluble uranium concentration was determined in the filtrate obtained by passing a 115 subsample through a centrifugal filter (30k molecular-weight-cut-off, Nanosep, Pall Life Sciences, 116 estimated 12 nm pore size). The U concentration was measured by LSC as described above. 117 After 10 days, half of the uranium-boehmite suspensions were amended with a small volume of a 0.05 M HEDPA stock solution (pH 7) to yield a suspension containing 5.4 mM HEDPA, 600 mg  $L^{-1}$ 118 119 boehmite, and 1M NaCl. The other half of the suspensions were amended with a 0.5 M HEDPA solution (pH 7) to yield a 50 mM, 600 mg L<sup>-1</sup>, 1 M NaCl suspension. The final U(VI) concentration was 5.8 µM in 120 121 all suspensions. Boehmite-free control solutions were also amended to 5.4 mM HEDPA as described 122 above. The soluble uranium concentration was determined via filtration followed by LSC as described 123 above. 124

# 125 **3. Results and Discussion**

# 126 **3.1 HEDPA Interactions with Boehmite: Sorption and Dissolution**

In the absence of complexants, the aqueous phase concentration of aluminum is limited by the formationof sparsely soluble Al (hydr)oxide solids, especially at circum-neutral pH values. As shown in Figure 2,

- 129 addition of HEDPA clearly enhances the dissolution of boehmite. Across the pH range 5 to 9, samples
- amended with HEDPA have aluminum concentrations significantly higher than the HEDPA-free control
- 131 system. The enhanced dissolution of boehmite in the presence of HEDPA is likely the results of formation

132 of  $Al_m H_h L_l$  complexes (where L represents completely deprotonated HEDPA and m, h, and l are 133 stoichiometric coefficients). The exact identity of specific complexes is uncertain at this point as the data 134 available in the current literature are not in agreement (5,6) and our potentiometric experiments to obtain 135 such data were unsuccessful due to the formation of a white precipitate across the neutral pH range (see 136 Section 2.3). The nature and composition of the white precipitate was not fully determined, but analysis 137 of the precipitate isolated from 1.0 M NaCl solutions with various Al(III):HEDPA molar ratios (from 1:1 138 to 1:2.5) indicated that it always had a 1:1 Al(III):HEDPA molar ratio. The observation of a 1:1 139 Al(III):HEDPA precipitate suggests that the low aluminum concentration across the neutral pH range 140 (Figure 2) may be controlled by the solubility of an Al(III):HEDPA solid phase. It was worth noting that 141 the constitution and concentration of the background electrolyte significantly affected the physical 142 characteristics of the precipitate. Precipitates prepared in 1.0 M NaCl and 1.0 M NaNO<sub>3</sub> tended to form 143 clearly distinct particles while precipitates from 1.0 M (CH<sub>3</sub>)<sub>4</sub>NCl and 1.0 M KCl tended to have a gel-144 like character, suggesting that sodium may cause significant aggregation of the Al(III):HEDPA particles. 145 In addition, the concentration of NaCl was also found to have a significant effect on the precipitate. 146 Solutions containing aluminum and HEDPA in a 2:5 molar ratio in ultra pure H<sub>2</sub>O, 0.1 M NaCl, or 1.0 M 147 NaCl were passed through filters with pore sizes ranging from 450 nm to 12 nm. The fraction of soluble 148 aluminum was found to be inversely related to the ionic strength of the solution. 149 The concentrations of aluminum in more acidic (pH 4 to 5) and basic (pH > 10) regions were

higher than those in the neutral pH region (Figure 2). After 2 days at pH 4 to 5, the concentrations of aluminum were comparable in systems with or without HEDPA. This is due to the formation of  $Al(OH)^{2+}$ and  $Al(OH)_{2^{+}}$  species yielding macromolar concentrations of aluminum in the absence of HEDPA. At pH > 10, the dominant aqueous species is  $Al(OH)_{4^{-}}$ , yielding high aluminum concentrations in systems

154 without HEDPA. At pH 11, the presence of HEDPA resulted in a lower aluminum concentration than that

155 in the control system up to 16 days (Figure 2). This difference may be due to precipitation of an

156 Al(III):HEDPA solid as described above or slow kinetics of dissociation of Al(III):HEDPA complexes

157 from the mineral surface. Data in the literature indicate that phosphate may form multi-nuclear surface

158 complexes which could inhibit detachment of the metal-ligand complex from the surface or even inhibit

159 dissolution (13). Therefore, as HEDPA surface complexation would be expected to occur through the

160 phosphate group, the inhibition of boehmite dissolution may be due to formation of a multi-dentate

161 surface complex.

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#### 163 **3.2 Sorption of U(VI) to synthetic boehmite in the absence of HEDPA**

164 Data describing the sorption of uranium in the absence of HEDPA are shown in Figure 3. In the absence 165 of HEDPA, approximately 15% of the uranium was sorbed at pH 4. As the pH increased, the fraction of 166 uranium sorbed increased until relatively steady partitioning was reached at pH > 5. Above pH 5, 167 approximately 90% of the total uranium was sorbed. The sorption edge occurring between pH 4 and 5 has 168 been preciously observed by a number of researchers examining uranium sorption to aluminum 169 (hydr)oxides (10, 14-15) This behavior is consistent with electrostatic attraction/repulsion between 170 uranium and the boehmite surface. As shown in Figure 1, the boehmite surface charge transitions from a 171 net positive to a net negative charge as the pH increases. This is due to the protonation and deprotonation 172 of aluminol surface sites. At low pH values the positively charged uranyl dioxycation is repelled by the 173 positively charged boehmite surface. As the pH was increased, both the net positive surface charge and 174 the positive charge on the U(VI) species decreased (the latter due to hydrolysis), resulting in stronger 175 sorption of U(VI) onto boehmite. The slight decrease in the fraction of uranium sorbed observed at higher 176 pH values is presumed to be due to formation of soluble uranyl-carbonates. Although experiments were 177 run in sealed centrifuge tubes, no effort was made to exclude carbonate from these experiments.

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#### 179 **3.3 Effect of HEDPA Concentration on U(VI) sorption to Boehmite**

180 Significant leaching of uranium from boehmite occurred rapidly following the amendment of the

181 equilibrated uranium-boehmite suspensions to either 5 mM HEDPA or 50 mM HEDPA (Figure 4a and

- 4b). At all pH values above 4.5, the addition of HEDPA resulted in rapid leaching of uranium from
- 183 boehmite. Four hours after amendment of the uranyl-boehmite suspensions with 5 mM HEDPA, the

184 fraction of uranium sorbed within the neutral pH region dropped from approximately 90% to 185 approximately 50% (Figure 4a). Over the next 181 days, the concentration of uranium in the aqueous 186 phase continued to increase. This slow step could be due to a rate limitation of the dissociation of a 187 U(VI):HEDPA complex from the boehmite surface or due to continued dissolution of the boehmite phase. 188 A similar effect was observed for the 50 mM HEDPA system although a larger fraction of the total 189 uranium was leached into the aqueous phase within the first 4 hours (Figure 4b). Furthermore, the 190 concentration of uranium in the aqueous phase after 181 days was significantly greater in the 50 mM 191 HEDPA suspension relative to the 5 mM HEDPA suspension. The increased partitioning of uranium to 192 the aqueous phase in the 50 mM HEDPA system is presumably due to enhanced dissolution of the 193 boehmite solid phase, saturation of the remaining boehmite surface sites with HEDPA, and/or formation 194 of U(VI):HEDPA complexes. As shown in Figure 2, after 135 days, at least 10% of the aluminum in 195 boehmite suspensions with 50 mM HEDPA is soluble at all pH values examined (Al concentration greater 196 than  $1.0 \times 10^{-3}$  M). At pH values less than 5, the boehmite was almost completely dissolved. Therefore the 197 total boehmite surface available for sorption of uranium or U(VI):HEDPA complexes was significantly 198 diminished in the 50 mM HEDPA system.

199 Data in Figure 4a (for the 5 mM HEDPA system) indicate a loss of uranium from the aqueous 200 phase at low pH values following the addition of HEDPA. Such loss appears to be consistent with the 201 precipitation of U(VI):HEDPA complexes that was observed at low pH and low HEDPA:U(VI) ratios 202 (<2) by Reed et al. (4). However, precipitation of a U(VI):HEDPA complex in these sorption experiments 203 is not expected, because the HEDPA concentration is between 3 and 4 orders of magnitude higher than 204 the uranium concentration and uranium was found to remain in the aqueous phase in all control (with 205 HEDPA but without boehmite) solutions spanning the pH range 4 to 11 for the duration of these 206 experiments. Possible reasons for the observed loss of U(VI) in the low pH regions of the 5 mM HEDPA 207 system are discussed in Section 3.4.

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#### 209 3.4 Effect of pH and HEDPA on U(VI) sorption to Boehmite

210 3.4.1 Uranium partitioning in the acidic pH region: Unless specified, the effects of pH on uranium 211 partitioning will be discussed below in terms of the 5.4 mM HEDPA system (Figure 4a). At pH 4, the 212 addition of HEDPA increased the fraction of uranium sorbed from 15% to 47% within 4 hours (the 213 fraction remained unchanged for the remainder of the experiment up to 181 days). This is the only sample 214 in which the addition of HEDPA caused an increase in the sorption of uranium. The uranium partitioning 215 at low pH observed in this system is similar to the observations of Cheng et al. (16) who found that 216 phosphate caused increased uranium sorption to goethite-coated sands at low pH values, relative to 217 phosphate free suspensions. A similar pH effect was also observed by Guo et al. (17) and Guo et al., (18) 218 when examining thorium and uranium sorption in alumina suspensions amended with phosphate. Guo et 219 al., (18) proposed that the enhanced sorption of uranium was due to formation of ternary surface 220 complexes with U(VI) and phosphate.

221 Assuming that the behavior of the phosphate groups of HEDPA may be similar to that of 222 phosphate, the increased uranium sorption observed in this work was also likely caused by the formation 223 of a ternary U-HEDPA-AlOH surface complex. As the speciation plot of U(VI) in a 5.4 mM HEDPA 224 solution shows, a number of anionic U(VI):HEDPA species exist in low pH regions (Figure 5). On the 225 other hand, the boehmite surface carries a net positive charge at low pH values. Therefore the observed increase in sorption could be due to sorption of anionic species such as  $UO_2L_2H_4^{2-}$  or  $UO_2L_2H_3^{3-}$ . Similar 226 227 ternary surface complexes have been proposed to explain increased metal sorption at low pH values in the 228 presence of anionic metal-ligand complexes (19-21). In this case, sorption of the metal is proposed to be 229 due to the formation of a ternary complex where the ligand bridges the mineral surface and the metal. 230 Such a geometry could be conceptualized utilizing the two phosphate groups of HEDPA. Within this pH region (pH ~4), HEDPA will be present as  $H_2L^{2-}$  (based upon the pK<sub>a</sub> values reported by Reed et al., (4)) 231 232 and will be attracted to the positively charged boehmite surface. Nowack and Stone (22) examined 233 HEDPA sorption to goethite and found that at low pH values, deprotonated anionic HEDPA species were 234 attracted to the positively charged goethite surface. As the pH increased, and the surface developed a net

negative charge, the sorption of HEDPA decreased (22). He et al., (23) observed similar pH effects when
examining phosphate sorption to alumina.

237 3.4.2 Uranium partitioning at circumneutral pH : Across the pH region from 5 to 8, the fraction 238 of sorbed uranium decreased significantly following the addition of HEDPA (Figure 4a). After 1 and 6 239 days, the fraction of uranium sorbed increased as the pH was increased from approximately 5 to 6.5. 240 Across the pH range 6.5 to 9, a relatively steady partitioning of uranium was observed for the sampling 241 intervals at 0.2, 1, and 6 days. After 6 days, a more general trend appeared in the data where the fraction 242 of uranium sorbed was found to increase as the pH increased from pH 5. It is noteworthy that 243 coprecipitation of uranium with an Al(III):HEDPA complex as discussed above is also a possibility. 244 However, the increase in uranium sorption with increasing pH is not consistent with coprecipitation as the 245 primary mechanism for loss of uranium from the aqueous phase. Across this circumneutral pH region (5 to 9), the anionic U(VI):HEDPA species  $UO_2L_2H_2^{4-}$ ,  $UO_2L_2H^{5-}$ , and  $UO_2L_2^{6-}$  are predominant (Figure 5) 246 247 and the boehmite surface maintains a net positive surface charge (Figure 1). This will allow for an 248 attractive force between the anionic U(VI):HEDPA complexes and the positively charged sites on the 249 boehmite surface. Since both free HEDPA and free U(VI) are known to sorb within this pH region, a 250 ternary surface complex could form through either uranium or HEDPA.

251 3.4.3 Uranium partitioning above pH 9: Above pH 9, the fraction of sorbed uranium decreased 252 significantly following the addition of HEDPA (Figure 4a). There was a slight decrease in the fraction of 253 uranium sorbed over the first 59 days followed by a significant decrease after 180 days. The rate at which 254 uranium was leached into the aqueous phase was similar to the rate of boehmite dissolution (Figure 2), 255 indicating that dissolution was a primary factor controlling uranium partitioning within this pH region. At 256 earlier time intervals, the fraction of uranium sorbed across the pH region 9 to 11 was higher relative to 257 the circumneutral pH region. This difference is proposed to be due to a change in the partitioning of 258 HEDPA within this pH region. Nowack and Stone (22) observed a significant decrease in the fraction of 259 HEDPA sorbed to goethite above pH 9. This is consistent with the full deprotonation of HEDPA in 1.0 M 260 NaCl ( $pK_{a4} = 9.5$ , measured in this work, data not shown). Therefore, boehmite sorption sites which are

occupied by HEDPA at neutral and acidic pH regions may be available for sorption of uranyl either as a
uranyl hydroxide or as a U(VI):HEDPA complex. In the absence of a complexing ligand, uranium
strongly sorbs to aluminum (oxy)hydroxides at high pH values (Figure 3 and references 10, 14, 15).
Therefore formation of a U(VI):hydroxide or U(VI):HEDPA (whereby surface complexation would occur
through uranium rather than through the phosphate groups of HEDPA) surface complex is feasible.
However, verification of such complexes through spectroscopy would be desirable, if not necessary, prior
to incorporation of such species into modeling efforts.

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# 269 4. Conclusions

270 Results from this work show that HEDPA is capable of significantly enhancing the solubility of 271 aluminum hydroxides. The degree of enhancement is dependent upon the bulk solution pH and the 272 concentration of HEDPA. HEDPA is capable of leaching uranium from synthetic boehmite through solid 273 phase dissolution and/or the formation of U(VI):HEDPA complexes. Across the pH range examined, the 274 speciation of U(VI):HEDPA complexes affects the partitioning of uranium between the solid and aqueous 275 phase. Partitioning of uranium was observed to vary with pH and correlated with the partitioning of 276 HEDPA and HEDPA-promoted boehmite dissolution. The sorption of uranium to boehmite in the 277 presence of HEDPA is proposed to be through ternary U(VI):HEDPA surface complexes. However, no 278 spectroscopic evidence is provided to indicate formation of such complexes. These results indicate that 279 HEDPA could be used to reduce the volume of the aluminum component of sludge within the Hanford 280 waste tanks. However, because a significant fraction of uranium (and presumably other actinides) could 281 also be leached from boehmite by the addition of HEDPA, careful consideration of the partitioning of the 282 actinides must be made if a strong complexant such as HEDPA is selected for sludge washing.

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348 349 Figure 1: Potentiometric titrations of 100 g  $L^{-1}$  boehmite suspensions in 0.01 M NaCl ( $\diamond$ ), 0.10

M NaCl ( $\Box$ ), and 1.0 M NaCl ( $\triangle$ ). 350





Figure 2: Effect of HEDPA on synthetic boehmite ( $\gamma$ -AlOOH) dissolution, [ $\gamma$ -AlOOH] = 600 mg 354 L<sup>-1</sup> (10 mM as Al<sup>3+</sup>), [NaCl] = 1.0 M. (A) [HEDPA] = 5.4 mM; (B) [HEDPA] = 50 mM.

Symbols: ( $\diamond$ ) 2 days, ( $\Box$ ) 16 days, ( $\blacktriangle$ ) 36 days, ( $\diamond$ ) 86 days, and ( $\Box$ ) 135 days. ( $\star$ ) Control system

356 contains no HEDPA and was measured after 135 days. Error bars, typically contained within

area of symbol at 95% certainty, have been removed for clarity.





359 Figure 3: Fraction of uranium sorbed to boehmite versus pH after 10 day equilibration. System parameters:  $[\gamma$ -AlOOH] = 0.66 g L<sup>-1</sup>;  $[U(VI)] = 6.1 \mu$ M; [NaCl] = 1.0 M.



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Figure 4: Effect of 5 mM (A) or 50 mM HEDPA (B) on Uranium Sorption to Boehmite:

Symbols ( $\blacktriangle$ ) 0.2 days, ( $\blacklozenge$ ) 1 day, ( $\vartriangle$ ) 6 days, ( $\blacksquare$ ) 14 days, ( $\blacksquare$ ) 30 days, ( $\square$ ) 59 days, and ( $\blacklozenge$ ) 180 days. For comparison, data from Figure 3 showing steady state distribution (10 day equilibrium) of U(VI) without HEDPA present is shown (×). System parameters: [ $\gamma$ -AlOOH] = 600 mg L<sup>-1</sup>; [NaCl] = 1.0 M; [U(VI)] = 5.8  $\mu$ M. Error bars removed for clarity, average 2 $\sigma$ = 3% propagated from counting statistics.



369 Figure 5: Speciation of U(VI) in 5.4 mM HEDPA and 1.0 M NaCl, calculated with the thermodynamic constants reported by Reed et al., (4). L stands for the fully deprotonated

HEDPA anion.