

1	The influence of calcium, sodium and bicarbonate on the uptake of uranium onto nanoscale
2	zero-valent iron particles
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24 Abstract

25 This work investigates the influence of calcium (Ca), sodium (Na) and bicarbonate (HCO_3^{-}) on the uptake of uranium (U) onto nanoscale zero-valent iron particles (nZVI). Batch systems 26 27 containing U at 1 mg/L and HCO₃⁻ at 0, 10, 100 and 1000 mg/L were tested with nZVI at 500 28 mg/L. NaCl was also added to the batch systems containing HCO₃⁻ at 0, 10 and 100 mg/L in 29 order to normalise the ionic strength to the batch system containing HCO_3^- at 1000 mg/L. Comparator systems were tested which contained U at 1 mg/L, HCO₃⁻ at 0, 10 and 100 mg/L 30 31 and equal moles of CaCl₂ to NaCl in the aforementioned batch systems. Mine water 32 containing a similar concentration of U (1.03 mg/L) was also tested as a natural analogue. Results demonstrate Ca as having no appreciable influence on the capacity of nZVI for U 33 34 uptake, with >99 % removal recorded for all systems. U desorption was enhanced, however, 35 with 87.3, 85.2 and 84.7 % removal recorded after 672 hours for the 0, 10 and 100 mg/L 36 bicarbonate systems compared with 99.9, 99.7 and 97.1 % recorded for the Na-bearing (Ca 37 absent) systems. Moreover, maximum U removal onto nZVI was directly proportional to 38 HCO₃⁻ concentration for the Na-bearing systems, whereas no trend was identified for the Ca-39 bearing systems. Results demonstrate Ca as having a significant inhibitive influence on the long-term retention (e.g. > 48 hours) of U on nZVI, which is independent of HCO_3^{-1} 40 41 concentration when it is also present at <100 mg/L.

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Keywords: uranium, nanoscale zero-valent iron particles, bicarbonate, calcium, mine water,
remediation

49 Introduction

50 A key environmental legacy of mankind's military and civil nuclear activities has been the 51 release of uranium (U) into the environment. U presents a considerable long-term 52 environmental concern and can significantly limit the potential for site redevelopment. In addition, the contamination of groundwater by more soluble forms of U can compromise 53 54 drinking water sources and spread contamination over significant distances. U mobility in groundwater is governed by its redox, sorption and complexation behaviour. It can exist in 55 56 many oxidation states [e.g. U(0), U(III), U(IV), U(V) and U(VI)], however, in natural groundwater systems (e.g. 6 < pH < 9) U(VI) it typically predominates as the uranyl ion 57 $(UO_2^{2^+})$, which is prone to complexation with many ubiquitous groundwater constituents, 58 59 including: phosphate, silicate, sulphate, fluoride and (bi)carbonate [1]. Consequently, more 60 than 42 dissolved U species, 89 U minerals and 368 inorganic crystal structures that contain 61 U(VI) have been documented [2]. The affinity of U with bicarbonate is of great importance due to the extremely high stability of uranyl carbonate complexes in comparison to the other 62 63 complexes which form in ground water systems [1]. In the presence of bicarbonate (HCO_3^{-}) and dissolved calcium (Ca), two ternary aqueous uranyl complexes: Ca₂UO₂(CO₃)₃ (log β_{213}) 64 = 30.70) and CaUO₂(CO₃)₃²⁻ (log β_{113} = 27.18) become increasingly prominent, and are 65 typically the dominant species in natural groundwater conditions [3]. For example, analysis 66 67 of groundwater samples taken from a U mill tailing remedial action (UMTRA) site in Tuba 68 City, Arizona, USA, determined that they comprised >99% of the U(VI) in solution [4]. In 69 addition, Fox et al. (2006) [5] recorded a decrease from 77% U(VI) adsorbed on quartz with no dissolved Ca to 10% adsorbed with a Ca concentration of 8.9 mM in carbonate-bearing 70 71 solution. As a consequence there is a great need for further studies investigating the 72 geochemistry of uranyl-calcium-carbonato complexes, in order to inform the development of 73 new water treatment technologies. In recent years much focus has been applied on the

74 potential utility of nanoscale zero-valent iron particles (nZVI) for the removal of aqueous U 75 from waste waters [6], [7], [8], [9], [10], [11], [12], [13], [14], [15], [16], [17]. Specific 76 mechanistic and kinetic investigations into the removal of uranyl-calcium-carbonato complexes onto nZVI, however, are limited at present [18]. Yan et al., (2010) [17] 77 78 investigated the uptake of U onto nZVI in anoxic conditions batch systems containing 79 NaHCO₃ with and without Ca. The work demonstrated that the kinetics of aqueous U(VI) removal is influenced by both Ca and HCO3⁻ concentration, namely that the kinetics of 80 81 aqueous U(VI) removal in mixed systems (1 mM HCO3⁻ and Ca) is comparable to those in 82 systems containing 1mM HCO₃⁻ or Ca. Overall, the results demonstrate nZVI as highly effective for the removal of U(VI) from anoxic waters for Ca and HCO₃⁻ concentrations up to 83 84 1 mM and 10 mM respectively. To the best of our knowledge, however, the long-term (e.g. 85 >96 hours) retention of U on nZVI in the presence of HCO₃⁻ and/or Ca is yet to be 86 investigated. The objective of this work is to investigate this process in batch systems also 87 containing dissolved oxygen (DO). The work has been established in order to correlate U 88 desorption processes with changes in nZVI corrosion production formation, in order to 89 simulate the eventual recovery of a nZVI subsurface treatment zone to redox conditions prior 90 to nZVI injection. Results are intended to evaluate the long-term integrity of in situ U 91 treatment using nZVI, namely the potential for U desorption following the ultimate exposure 92 of the treatment zone to DO.

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94 2. Materials and methods

95 **2.1. Solution synthesis**

All chemicals used in this study [CaCl₂, C₂H₆O, C₃H₆O, FeSO₄·7H₂O, HNO₃, NaCl, NaBH₄,
NaHCO₃, NaOH and (UO₂(CH₃COO)₂·2H₂O)] were of ACS reagent grade and all solutions
were prepared using Milli-Q purified water (resistivity >18.2 MΩ cm). Four 400 mL stock

99 solutions were synthesised comprising U at 1 mg/L and HCO₃⁻ at 0, 10, 100 and 1000 mg/L 100 using (UO₂(CH₃COO)₂·2H₂O) and NaHCO₃ respectively. NaCl was then added at 696, 688 101 and 625 mg/L to the batch systems containing 0, 10 and 100 mg/L NaHCO₃ respectively in 102 order to equalise their ionic strength with the batch system containing 1000 mg/L NaHCO₃. 103 A further three 400 mL solutions were synthesised comprising U at 1 mg/L, NaHCO₃ at 0, 10 104 and 100 mg/L. Equal moles of Ca to Na that was added to the aforementioned 0, 10 and 100 105 mg/L NaHCO₃ solutions, was also added to the batch systems, which comprised 719, 712 and 106 647 mg/L of CaCl₂ respectively. The pH of all systems was then adjusted to 8.0 using 0.5 M 107 NaOH. The solutions were stored in sealed glass jars (500 ml Schott Duran) in the open 108 laboratory prior to analysis and in between sampling intervals.

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110 **2.2. U-bearing mine water**

111 The U-bearing natural water used herein was taken from the Ciudanovita Uranium Mine, 112 Banat, Romania. The site is valley confined and bounded by limestone ridges which 113 contribute significant concentrations of dissolved HCO_3^- to ground and surface waters. The 114 water is used for mining purposes and is pumped from approximately 200 m below sea level. 115 It initially contains trace concentrations of DO, however, quickly equilibrates with the 116 atmosphere to reach DO concentrations more typical for that of vadose and/or surface waters 117 (7-12 mg/L), changing its redox potential and associated U(VI) transport properties in the 118 process.

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120 **2.3. Zero-valent iron nanoparticle synthesis**

Pure nZVI were synthesised following the method first described by Glavee et al., (1995) [12] [19], and then adapted by Wang and Zhang (1997) [20], which uses sodium borohydride to reduce ferrous iron to a metallic state. 7.65 g of FeSO4·7H₂O was dissolved in 50 mL of 124 Milli-Q water (> 18.2 M Ω cm) and then a 4M NaOH solution was used to adjust the pH to 6.8. NaOH addition was performed slowly, dropwise, to avoid the formation of 125 hydroxocarbonyl complexes. The salts were reduced to nZVI by the addition of 3.0 g of 126 127 NaBH₄. The nanoparticle product was isolated via centrifugation (Hamilton Bell v6500 Vanguard centrifuge, 6500 RPM for 2 minutes), rinsed with Milli-Q water (ratio of 50 mL 128 129 per g of nZVI) and then centrifuged (Hamilton Bell v6500 Vanguard centrifuge, 6500 RPM for 2 minutes). This step was then repeated but using ethanol and then acetone as the solvent. 130 The nanoparticles were dried in a vacuum dessicator (approx. 10⁻² mbar) for 48 hours and 131 132 then stored in an argon filled (BOC, 99.998 %) MBraun glovebox until required.

133

134 **2.4. Experimental procedure**

135 Eight 500 mL Schott Duran jars were each filled with 400 mL of each U-bearing solution. A 136 nZVI mass of 0.2 g (0.5 g/L) was then added. In each case, the nZVI was suspended in 5 mL 137 of ethanol and dispersed by sonication for 30 seconds. Each batch system was sampled at 0 h, 138 0.5h, 1 h, 2 h, 24 h, 48 h, 72 h, 7 d, 14 d and 28 d. Prior to sampling, the jars were gently 139 agitated to ensure homogeneity and pH, Eh and DO measurements were taken using a Hach 140 multimeter (model HQ40d) using a combination gel electrode for pH measurements, a gel-141 electrolyte reference ORP electrode for Eh measurements, and a luminescent/optical 142 dissolved oxygen (LDO) probe for DO measurements. Aliquots of 5 mL were then taken 143 from each batch system and centrifuged using a Hamilton Bell Vanguard V6500 desktop centrifuge at 6500 rpm for 30 seconds to separate the liquid and solid phases. The supernatant 144 was then decanted, filtered through a 0.22 µm cellulose acetate filter and then prepared for 145 146 inductively coupled optical emission spectrometry (ICP-OES) and inductively coupled 147 plasma mass spectrometry (ICP-MS) by the additional of 1 % by volume of concentrated 148 HNO₃. The solid was prepared for X-day diffraction (XRD) by rinsing with Milli-Q water (ratio of 50 mL per g of nZVI) and then centrifugation (Hamilton Bell v6500 Vanguard centrifuge, 6500 RPM for 2 minutes). This step was then repeated but using ethanol and then acetone as the solvent. The resultant suspension was then pipetted onto a glass optical microscope slide and dried in a vacuum chamber at $<1 \times 10^{-5}$ mbar for 2 hours prior to XRD analysis.

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155 **2.4. Sample analysis methods**

156 **2.4.1. BET surface area analysis**

157 In preparation for BET surface area analysis, samples were degassed under vacuum $(1 \times 10^{-2} \text{ mbar})$ for a 24 hour period at a temperature of 75°C. A known mass of the dried material was 159 then measured using a Quantachrome NOVA 1200 surface area analyser, with N₂ as the 160 adsorbent and following a 7 point BET method.

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162 2.4.2. ICP-AES preparation and conditions

Blanks and standards for ICP-AES analysis were prepared in 1 % nitric acid, with Fe standards of 0.10, 0.2, 0.3, 0.4 and 0.5 mg/L. An Agilent 710 ICP-OES (sequential spectrometer) fitted with a cyclone spray chamber and a teflon mist nebulizer was used. The Fe-concentration was measured using the emission line at 234.35 and 259.94 nm.

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168 2.4.3. ICP-MS preparation and conditions

Samples for ICP-MS analysis were prepared by performing a 20 times dilution in 1 % nitric
acid (analytical quality HNO₃ and Milli-Q water). Blanks and U standards at 1.0, 2.0, 10, 20

171 and 50 $\mu g/L$ were also prepared in 1 % nitric acid. The ICP-MS instrument used was a

172 Thermo Scientific PlasmaQuad 3.

2.4.4. Transmission electron microscopy 174 Transmission electron microscopy (TEM) images were obtained with a JEOL JEM 1200 EX 175 176 Mk 2 TEM, operating at 120 keV. Nanoparticle samples were suspended in absolute ethanol 177 via sonication for 120 seconds and then mounted on 200 mesh holey carbon coated copper grids. The nanoparticle infused grids were then dried in a vacuum chamber at $< 1 \times 10^{-5}$ mbar 178 179 for 2 hours. 180 181 2.4.5. X-ray diffraction 182 A Phillips Xpert Pro diffractometer with a $Cu_{K\alpha}$ radiation source (λ = 1.5406 Å) was used for XRD analysis (generator voltage of 40 keV; tube current of 30 mA). XRD spectra were 183 acquired between 20 angles of 10–90°, with a step size of 0.02° and a 2 s dwell time. 184 185 186 2.4.6. X-ray photoelectron spectroscopy 187 A Thermo Fisher Scientific Escascope equipped with a dual anode X-ray source (Al_{K α} 1486.6 eV and Mg_{Ka} 1253.6 eV) was used for XPS analysis. Samples were analysed at $<5 \times 10^{-8}$ mbar 188 with Al_{Ka} radiation of 300 W (15 kV, 20 mA) power. High resolution scans were acquired 189 190 using a 30 eV pass energy and 300 ms dwell times. Following the acquisition of survey 191 spectra over a wide binding energy range, the Fe 2p, C 1s, O 1s and U 4f spectral regions 192 were then scanned at a higher energy resolution such that valence state determinations could 193 be made for each element. Data analysis was carried out using Pisces software (Dayta 194 Systems Ltd.) with binding energy values of the recorded lines referenced to the adventitious hydrocarbon C1s peak at 284.8 eV. In order to determine the relative proportions of Fe²⁺ and 195 Fe^{3+} in the sample analysis volume, curve fitting of the recorded Fe 2p photoelectron peaks 196

197 was performed following the method of Grosvenor et al. (2004) [21]. The Fe 2p profile was 198 fitted using photoelectron peaks at 706.7, 709.1, 710.6 and 713.4 eV corresponding to Fe⁰, 199 $Fe^{2+}_{octahedral}$, $Fe^{3+}_{octahedral}$ and $Fe^{3+}_{tetrahedral}$. These parameters were selected on the basis that the 200 surface oxide was assumed to be a mixture of wüstite and magnetite, as the oxide Fe^{2+} is in 201 the same coordination with the surrounding oxygen atoms in both forms of oxide.

202 **3. Results and discussion**

3.1. Characterisation of the unreacted nanoparticles

204 Preliminary characterisation of the nZVI was performed using BET surface area analysis, 205 TEM, XRD and XPS. BET surface area analysis determined that the surface area of the nZVI 206 was 16.97 m²/g. TEM analysis determined that the nZVI were roughly spherical, with an 207 approximate size range of 20-120 nm and an average diameter of 32 nm. The density contrast between the Fe⁰ core and oxide shell within the nZVI was not identified and the material was 208 209 recorded as relatively amorphous. Individual particles were recorded as aggregated into 210 chains and rings, attributed to electrostatic and magnetic attraction forces between adjacent particles. XRD analysis recorded a broad diffraction peak at 44.9° 20 and two lower intensity 211 peaks at 65.6 and 82.6 20, confirming the presence of amorphous Fe⁰. XPS analysis recorded 212 the outer surface of the nZVI to be comprised of a mixed valent (Fe^{2+}/Fe^{3+}) oxide. The results 213 214 are summarised in Table 1.

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Parameter		nZVI
	0-50 nm	85
Particle size distribution (%)	50-100 nm	8
	>100 nm	7
Crystallinity		Amorphous (α-Fe)

Surface area (m^2/g)		16.97
	Fe	30.5
Surface composition (%)	0	32.1
	С	14.5
	В	22.9
	$(Fe^{0}/Fe^{2+} + Fe^{3+})$	0.02
Surface stoichiometry	Fe ²⁺ /Fe ³⁺	0.38

Table 1. A summary of the experimental results regarding the bulk and surface properties of
the nZVI used in the current work. Note: a significant proportion of the carbon detected is
likely to be adventitious carbon.



Figure 1. TEM images of the unreacted nZVI.



Example 223 Binding Energy (eV) 224 Figure 2. XRD spectra for the range of 10-90° 2 θ (LHS) and curve fitted XPS Fe 2p_{3/2} 225 photoelectron spectra (RHS) for the unreacted nZVI.

227 **3.2.** Preliminary characterisation of the U contaminated water

Prior to nanoparticle addition, the U-bearing mine water was characterised using ICP-MS (U), ICP-AES (Ca, Mg, Na and U), volumetric titration (HCO_3^-) and ion chromatography (NO_3^- and SO_4^{2-}), with the results displayed in Table 2.

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Chemical species	Concentration (mg/L)
Major cations	
Ca	52
Mg	7
Na	652
U	1.03
Major anions	
HCO ₃ -	845
NO ₃ -	0.9

SO4 ²⁻	65

Table 2. Concentrations of U and major ions present in the mine water, analysed by ICP-MS (U), ICP-AES (Ca, Mg, N and U), volumetric titration (HCO_3^-) and ion chromatography (NO_3^- and SO_4^{2-})

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237 **3.3. Changes in pH/Eh/DO**

238 Following the addition of the nZVI a rapid shift to strongly chemically reducing conditions 239 was recorded, with Eh less than -350 mV and near-zero DO (<0.5 mg/L) recorded for all 240 batch systems after 30 minutes reaction time (Figure 3). An accompanying increase in pH 241 was also recorded for all systems (Figure 3) with the behaviour therefore attributed to the rapid oxidation of nanoparticulate surfaces, consuming DO and H⁺ and increasing the 242 243 reduction potential of the system (Eq. 1-4). In the early stages of reaction (<30 minutes), the predominant mechanism of nZVI corrosion is considered to have been through reaction with 244 H⁺ via the consumption of DO (Eq. 1 and 2). Following this time period the absence of DO 245 246 dictates that corrosion could only proceed through the direct reaction (hydrolysis) with water (Eq. 3 and 4). Whilst Fe⁰ is an essential component of these reactions it is noted that the nZVI 247 248 used in the current work was determined as having a encapsulating (hydr)oxide layer which was acquired during synthesis (Figures 1 and 2). Fe^{0} is therefore considered to only react 249 250 indirectly with the aqueous solutions and associated dissolved components.

251

252
$$2Fe^{0}_{(s)} + 4H^{+}_{(aq)} + O_{2(aq)} \rightarrow 2Fe^{2+} + 2H_2O_{(l)}$$
 $E^{0} = +1.67 \text{ V}$ (Eq. 1)

253
$$2Fe^{2+}(s) + 2H^{+}(aq) + \frac{1}{2}O_{2}(aq) \rightarrow 2Fe^{3+} + H_2O_{(l)}$$
 $E^0 = +0.46 \text{ V}$ (Eq. 2)

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$$\operatorname{Fe}^{0}_{(s)} + 2H_{2}O_{(l)} \rightarrow \operatorname{Fe}^{2+} + H_{2(g)} + 2OH_{(aq)}^{-} \qquad E^{0} = -0.39 \text{ V}$$
 (Eq. 3)

255 $2Fe^{2+}{}_{(s)} + 2H_2O_{(l)} \rightarrow 2Fe^{3+} + H_{2(g)} + 2OH^{-}{}_{(aq)} \qquad E^0 = -1.60 \text{ V}$ (Eq. 4)

256 For all systems chemically reducing Eh and low DO (<3 mg/L) were maintained for the first 257 24 hours of reaction, with a gradual reversion to ambient conditions recorded onwards from this time. A concurrent decrease in solution pH was also recorded for all systems with the 258 259 behaviour therefore ascribed to a decrease in the rate of nZVI oxidation (corrosion). No clear trend in Eh, DO or pH change as a function of HCO₃⁻ concentration can be discerned, with 260 261 the behaviour recorded to be relatively similar for all systems. This was expected, given that 262 each synthetic solution had an equal ionic strength, which is a key influence on nZVI 263 corrosion rate.

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Figure 3. Changes in solution Eh, dissolved oxygen (DO) and pH as a function of reaction time (0-672 hrs) for the batch systems containing nZVI. The control (nanoparticle free) for each batch system (not shown) recorded an Eh variation of <10 mV, a DO variation of <0.2 mg/L and a pH variation of <0.05 from the starting solution.

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271 **3.4.** Changes in aqueous U and Fe concentrations

272 Analysis of liquid samples using ICP-MS recorded rapid U_(aq) removal in all systems with 273 removal of >50% recorded after 30 minutes in all systems (Figure 4). It can be noted that relatively similar behaviour was recorded for the batch systems containing 0 and 10 mg/L 274 275 NaHCO₃ (no Ca), with 99.9 % U_(aq) removal recorded within the first 24 hours for both 276 systems. Slightly lower U uptake was recorded for the 100 mg/L NaHCO₃ (no Ca) batch 277 system, with maximum U removal of 98.8 % U removal recorded after 24 hours. In contrast a 278 maximum of only 62 % U uptake was recorded for the 1000 mg/L NaHCO₃ (no Ca) system, 279 with near total re-release occurring from 72 hours onwards. Comparing these results with U 280 uptake data recorded for the batch systems containing Ca it can be noted that for all systems containing Ca similarly high initial U uptake (> 99 %) was recorded in the first 24 hours of 281 282 reaction, however, desorption of 12.7, 14.8 and 15.3 % was recorded for the batch systems 283 containing 0, 10 and 100 mg/L NaHCO₃ (Figure 5). Similar behaviour was also recorded for 284 the U-bearing mine water, with >99.5 % uptake recorded after 30 minutes of reaction, but followed by near-total re-release in the latter stages of the reaction. Overall the results 285 286 demonstrate that Ca exhibits a strong control on the long-term stability of U on nZVI in 287 waters containing DO and at circumneutral pH.



Figure 4. Changes in aqueous U and Fe concentrations as a function of reaction time (0-672
hrs) for the batch systems containing nZVI. The control (nanoparticle free) for each batch

system (not shown) recorded an aqueous U concentration variation of <0.2 mg/L from the
starting solution.





Figure 5. Aqueous U concentration as a function of bicarbonate concentration (0 – 100 mg/L) for the batch systems containing Na (LHS) and Ca (RHS).

298 **3.5. X-ray diffraction**

299 XRD was used to determine the bulk crystallinity and composition of nZVI solids extracted 300 from all batch systems at periodic intervals during the sorption experiments (Figure 6). During the initial stages of the reaction (≤ 4 hours) a transition from Fe⁰, with peaks centred 301 at 44.6 and 82.6° 20 corresponding to Fe(110) and Fe(211), to a mixture of 2-line ferrihydrite 302 303 (5Fe₂O₃•9H₂O) and magnetite (Fe₃O₄) was recorded for all systems. This was not unexpected given that both 5Fe₂O₃•9H₂O and Fe₃O₄ are common Fe⁰ corrosion products in circumneutral 304 305 and alkaline solutions [22], [23], [24]. In general, 5Fe₂O₃•9H₂O was the most prevalent 306 corrosion product during the initial stages of nZVI corrosion (e.g. <1 day) with Fe₃O₄ 307 emerging as the most prevalent during the latter stages. This was expected, given that Fe₃O₄ 308 is known to readily form from 5Fe₂O₃•9H₂O in near-neutral to basic pH in the presence of 309 HCO3⁻. A mixture of 5Fe₂O₃•9H₂O and Fe₃O₄ was recorded after 28 days reaction for the 310 mine water and solutions containing NaHCO₃ and Na. In contrast, much greater intensity

- 311 diffraction peaks corresponding to Fe₃O₄ were recorded in the latter stages of the reaction (>
- 312 1 day) for the batch systems containing NaHCO₃ and Ca, indicating that Fe₃O₄ was present in
- 313 much greater quantity (relative to other corrosion products) and/or crystallinity.
- 314



Figure 6. X-ray diffraction spectra for the range of 10-90° 2θ for nZVI extracted from the
different solutions after 0h, 1h, 4h, 24h, 48h, 168h and 672h.

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319 **3.6. Implications for the** *in situ* treatment of aqueous U using nZVI

An intrinsic technical challenge associated with the *in situ* treatment of metal and metalloid contaminants (such as U) using nZVI is the prospect for contaminant remobilisation [11], [18]. Ca and HCO_3^- are ubiquitous groundwater constituents, and are also known to significantly enhance the aqueous mobility of U via the formation of uranyl-calciumcarbonato complexes [1]. Indeed it has been demonstrated in recent years that the vast majority of aqueous U present in circumneutral and alkaline pH natural waters is comprised 326 of uranyl-calcium-carbonato complexes, due to their significantly stronger binding affinity 327 compared to other common groundwater constituents [3]. This presents a considerable technical challenge for the treatment of U from natural waters, especially in situ treatment 328 329 applications, where U removal from the aqueous phase must be ensured for long-term (or even quasi-permanent) timescales. The work presented herein has key implications for the in 330 331 situ treatment of U using nZVI. It is suggested that because ultimate recovery in DO is a 332 seemingly unavoidable fate for a typical subsurface nZVI treatment zone, associated 333 corrosive transformation of nZVI into (hydr)oxides and a cessation of strongly chemically 334 reducing groundwater conditions (i.e. Eh < -400 mV), is likely to result in considerable U 335 desorption. Indeed considering the significant geochemical perturbation caused by nZVI 336 injection, subsurface treatment zones are often highly metastable, and even a gradual 337 reversion in groundwater conditions toward a pre-injection state may be enough for 338 significant U remobilisation to occur. It is therefore suggested that secondary processes 339 which would maintain strongly chemically reducing conditions (e.g. Eh < -400 mV) within 340 the treatment zone, such as periodic nZVI reinjection and/or capping the treatment zone using 341 an impermeable engineered layer, are essential in order to preserve the long-term 342 performance of nZVI for the in situ treatment of U.

343

344 **4.** Conclusions

This work presents the influence of Na, Ca and HCO_3^- on the removal of U onto nZVI at pH 8. Batch systems containing U at 1 mg/L and HCO_3^- at 0, 10, 100 and 1000 mg/L were tested with nZVI at 500 mg/L. NaCl was also added to the batch systems with HCO_3^- at 0, 10 and 100 mg/L in order to normalise the ionic strength. Comparator systems were tested which contained U at 1 mg/L, HCO_3^- at 0, 10 and 100 mg/L and equal moles of CaCl₂ to NaCl added to the aforementioned batch systems. Mine water of similar U concentration (1.03 351 mg/L) was also investigated as a natural analogue. Ca had no appreciable impact on the capacity of nZVI for U removal, with >99 % recorded for all systems; however, it 352 353 significantly enhanced U desorption, with 87.3, 85.2 and 84.7 % removal recorded after 672 354 hours for the 0, 10 and 100 mg/L HCO₃⁻ systems in comparison to 99.9, 99.7 and 97.1 % recorded for the system with Ca absent. Moreover, in systems were Ca was absent maximum 355 356 U removal onto nZVI was recorded as directly proportional to HCO₃⁻ concentration, whereas for systems containing Ca no clear trend was identified. Overall, the results demonstrate Ca 357 358 as significantly decreasing the long-term stability (e.g. > 48 hours) of U sorbed on nZVI in 359 the presence of DO, with U removal also independent of HCO_3^- concentration when it is also present at <100 mg/L. 360

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