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Transport and speciation of uranium in groundwater-surface water systems impacted by legacy milling operations

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20 ABSTRACT

21 Growing worldwide concern over uranium contamination of groundwater resources has placed 22 an emphasis on understanding uranium transport dynamics and potential toxicity in groundwater-23 surface water systems. In this study, we utilized novel in-situ sampling methods to establish the 24 location and magnitude of contaminated groundwater entry into a receiving surface water 25 environment, and to investigate the speciation and potential bioavailability of uranium in 26 groundwater and surface water. Streambed temperature mapping successfully identified the 27 location of groundwater entry to the Little Wind River, downgradient from the former Riverton 28 uranium mill site, Wyoming, USA. Diffusive equilibrium in thin-film (DET) samplers further 29 constrained the groundwater plume and established sediment pore water solute concentrations and 30 patterns. In this system, evidence is presented for attenuation of uranium-rich groundwater in the 31 shallow sediments where surface water and groundwater interaction occurs. Surface water grab 32 and DET sampling successfully detected an increase in river uranium concentrations where the groundwater plume enters the Little Wind River; however, concentrations remained below 33 34 environmental guideline levels. Uranium speciation was investigated using diffusive gradients in 35 thin-film (DGT) samplers and geochemical speciation modelling. Together, these investigations 36 indicate uranium may have limited bioavailability to organisms in the Little Wind River and, 37 possibly, in other similar sites in the western U.S.A. This could be due to ion competition effects 38 or the presence of non- or partially labile uranium complexes. Development of methods to establish 39 the location of contaminated (uranium) groundwater entry to surface water environments, and the 40 potential effects on ecosystems, is crucial to develop both site-specific and general conceptual 41 models of uranium behavior and potential toxicity in affected ground and surface water 42 environments.

44 **1. Introduction**

45 Uranium (U) is a radioactive element with a crustal concentration of approximately 2.7 mg kg⁻ ¹ (Gupta and Singh, 2003), and is found with concentrations of 12 ng L^{-1} to 4.8 µg L^{-1} in stream 46 water worldwide (Windom et al., 2000) and approximately 3.4 µg L⁻¹ in sea water (Dunk et al., 47 48 2002). It occurs in the environment as a result of natural weathering of U-rich sediments and rocks 49 or via anthropogenic activities such as mining, nuclear accidents and waste disposal, as well as 50 nuclear weapons testing and disposal (Abdelouas, 2006; Guo et al., 2016; Nolan and Weber, 2015). 51 In aqueous environments, U exists primarily in the hexavalent oxidation state (U(VI)) as the uranyl ion (UO₂²⁺) at pH <5, or as stable uranyl hydroxide or carbonate complexes (e.g. UO₂(CO₃) $_2^{2-}$) at 52 53 pH > 7 (Nolan and Weber, 2015). The tetravalent state (U(IV)) is much less soluble and known to 54 accumulate in anoxic sediments such as ore deposits and contaminated aquifers (Bone et al., 55 2017a; Bone et al., 2017b).

56 As a non-essential trace metal and radionuclide, U can be highly toxic due to both its chemical 57 (speciation) and radiological (isotopic composition) properties, and exposure to U through drinking water is associated with nephrotoxic effects (Bjørklund et al., 2020; Pinney et al., 2003). 58 59 As a consequence, the U.S. Environmental Protection Agency and the World Health Organization have set environmental guideline levels of 30 µg L⁻¹ in drinking water (U.S. Environmental 60 61 Protection Agency, 2000; World Health Organization (WHO), 2012). In recent years, U has 62 emerged as a widespread threat to human health and ecosystems due to elevated concentrations in 63 groundwater from historical uranium mining and processing, nuclear waste disposal, and natural 64 geogenic sources (Bone et al., 2017b; Coyte et al., 2018; Nolan and Weber, 2015). Furthermore, 65 the growing worldwide demand for groundwater as a source of water (Dalin et al., 2017), and for 66 U as a source of energy (World Nuclear Association, 2019), has raised concerns regarding the
67 future risks to humans and ecosystems of U-contaminated groundwater.

68 Uranium is one of the principal contaminants at former mill sites managed by the U.S. 69 Department of Energy's (DOE) Office of Legacy Management (Dam et al., 2015). Typically, U 70 ore was crushed and ground into small particles in the milling process before being leached to 71 dissolve U oxides into a concentrated liquid slurry. Decades of U ore processing for use in the U.S. 72 Government nuclear weapons and energy programs resulted in the accumulation of approximately 73 30 Mm³ of U ore processing waste (U.S. Department of Energy, 2020). Tailings that remained 74 after the milling process were generally held in unlined impoundments resulting in widespread 75 groundwater contamination in the western U.S.A. (U.S. GAO (U.S. Government Accountability 76 Office), 2020). Today, the environmental liability of the DOE is estimated at USD\$7.35 billion, 77 with approximately 40% of these costs associated with long-term monitoring and maintenance 78 (U.S. GAO (U.S. Government Accountability Office), 2020). Contamination of groundwater used 79 for drinking and irrigation water is a major concern (U.S. GAO (U.S. Government Accountability 80 Office), 2020). However, the interaction of U-rich groundwater with surface water systems and 81 the potential effects on freshwater ecosystems has not previously been considered. Yet, the ability 82 to locate groundwater plumes, to establish the effect on receiving surface water systems, and to 83 establish the potential bioavailability of U to freshwater ecosystems, are seen as key requirements 84 to meet the long-term monitoring and remediation objectives set out by the DOE (U.S. GAO (U.S. 85 Government Accountability Office), 2020).

Recognizing the challenges associated with monitoring the potential human and ecological
effects of U-contaminated groundwater, the DOE, in collaboration with the U.S. Geological
Survey, University of Montana, Liverpool John Moores University, and Northern Arapahoe Tribe,

conducted a pilot-scale investigation of groundwater-surface water interaction at the former U ore mill and processing site at Riverton, Wyoming, from 2015 to 2018. The overall aim of this research was to test and apply a range of hydrological, geophysical, ecological and biogeochemical tools and methods to identify and quantify the effect of groundwater U plumes on connected surface water environments. As such, the research presented in this contribution is part of a larger and ongoing interdisciplinary study – project data is available in Naftz et al. (2019) and Terry and Briggs (2019).

96 In this specific study, we present a novel approach to establish the water quality and potential 97 biological effects of contaminated (U) groundwater interaction with surface waters. The approach 98 utilizes streambed temperature mapping as an inexpensive and unobtrusive method for identifying 99 groundwater fluxes through the streambed at small spatial scales (Conant, 2004). Areas of 100 groundwater efflux are then targeted using diffusive equilibrium in thin-film (DET) samplers to 101 establish spatial and vertical patterns of U sediment pore water chemistry. Diffusive gradients in 102 thin-film (DGT) samplers, in conjunction with speciation modelling, are then used to investigate 103 the potential bioavailability of U in ground and surface waters. Both DET and DGT samplers have 104 previously been used to establish U concentrations and speciation in surface waters (Drozdzak et 105 al., 2016; Hutchins et al., 2012; Turner et al., 2014; Turner et al., 2012) and in sediments in 106 laboratory microcosm experiments (Gregusova and Docekal, 2013). However, our study is the first 107 to apply DET and DGT techniques in-situ to investigate U concentrations, transport and speciation 108 in a groundwater-surface water system. This represents a substantial contribution to our conceptual 109 understanding of U behavior and potential toxicity in affected ground and surface water 110 environments.

Our primary aim was to demonstrate the potential of using in-situ temperature, DET and DGT samplers, in conjunction with speciation modelling, to establish the effect of legacy U groundwater plumes on connected surface waters. The specific objectives were to: (1) establish the location, magnitude, and effect of contaminated groundwater efflux to surface water, and; (2) establish the speciation and potential bioavailability of U at the groundwater-surface water interface and in surface water systems.

117

118 **2. Methodological approach**

119 2.1 Study area

120 The Uranium Mill Tailings Radiation Control Act (UMTRCA) was enacted in the U.S.A. in 121 1978 to provide for the safe disposal and long-term stabilization of U mill tailings to minimize 122 environmental and human health risks (U.S. Department of Energy, 2020). Twenty-two mill sites 123 were effectively abandoned when UMTRCA was passed; these were designated as Title I sites 124 where DOE has responsibility for long-term monitoring and remediation, including removal of 125 residual surface wastes. Title II sites (six sites) continued to operate, or commenced processing 126 activities, after UMTRCA was passed, and responsibility for their remediation falls to the licensee. 127 The Riverton site is situated on an alluvial terrace in the Wind River Basin, Wyoming, U.S.A. 128 Groundwater flow direction is generally southeast and occurs in three aquifers beneath the site 129 (U.S. Department of Energy, 1998): (1) a shallow, unconfined aquifer, consisting of approximately 130 4.6 - 6 m of unconsolidated alluvial material, and underlain by a discontinuous shale confining 131 layer 2 - 3 m thick; (2) a middle, semi-confined aquifer; and (3) a deeper, confined aquifer 132 comprising the upper units of the Eocene Wind River Formation. The climate of the region is semi-133 arid to arid, with mean annual precipitation approximately 20 cm (the majority occurring April –

June as spring snows and showers) (U.S. Department of Energy, 1998). Phreatophytes cause evapotranspiration to vary daily and seasonally in response to seasonal climate and plant growth factors (Dam et al., 2015).

137 The Riverton Processing site (Title I) produced yellowcake (U_3O_8) , used in the preparation of 138 U fuel for nuclear reactors and weapons, from 1958 to 1963 and was located on the Wind River 139 Indian Reservation near Riverton, Wyoming (Figure 1). The mill processed a total of 816,470 140 tonnes of U ore that was mined in the Gas Hills mining district in Wyoming (Merritt, 1971). During 141 the milling process, ore was crushed and ground, and water was added to create a slurry. Mill 142 tailings that remained after extraction of U were conveyed by slurry to a 29 hectare unlined tailings 143 impoundment and stockpiled (U.S. Department of Energy, 1995). Following the UMTRCA, 144 surface remediation of the site was completed by DOE in November 1989. Approximately, 1.4 145 Mm³ of contaminated material was removed from the site; however, decades of leaching of the 146 tailings slurry has led to contamination of the shallow groundwater beneath and downgradient 147 from the site. The primary contaminant is U, but other trace elements (arsenic (As), boron (B), 148 iron (Fe), lead (Pb), manganese (Mn), mercury (Hg), molybdenum (Mo), nickel (Ni), selenium 149 (Se), vanadium (V)), and sulfate (SO₄) are above background concentrations (Narasimhan et al., 150 1986; U.S. Department of Energy, 1998; White et al., 1984).

151 The approach taken by DOE to the management of groundwater contamination at Riverton and 152 other UMTRCA sites was based on the natural flushing compliance strategy, where modelling of 153 contaminant movement in aquifers using the Groundwater Analysis and Network Design Tool 154 (GANDT) predicted U concentrations falling below the maximum contamination level (30 μ g L⁻ 155 ¹) within a 100 year regulatory timeframe (Dam et al., 2015). Prior to 2010, groundwater 156 monitoring indicated that U concentrations at the Riverton Processing site were decreasing at a 157 steady rate and were in general agreement with the GANDT modelling predictions (Dam et al., 158 2015; U.S. Department of Energy, 2009). However, a series of flood events beginning in June 159 2010 mobilized contaminants held in normally unsaturated materials above the alluvial water table, 160 as well as contaminants within the aquifer (Dam et al., 2015; Ranalli and Naftz, 2014), resulting 161 in substantial increases in groundwater U concentrations in some monitoring wells and possible 162 increased flux of contaminants to the Little Wind River. Groundwater monitoring data from other 163 U ore processing sites reveals similar diversions from conceptual models of U attenuation over 164 time (Shafer et al., 2014; Zachara et al., 2013). Further research on the interaction of legacy 165 groundwater plumes with surface waters is therefore needed to help refine natural flushing 166 predictions.

167

168 2.2 Streambed temperature mapping

169 A streambed temperature survey (following Conant, 2004) was conducted in August 2017 170 using a Traceable Control Company digital thermometer with a 10 cm probe. Measurement 171 locations were surveyed with sub-meter accuracy using a Trimble R1 GNSS Receiver. Streambed 172 temperature mapping areas were constrained to potential areas where the legacy groundwater 173 plume intersected the left bank of the Little Wind River. Mapped areas were further constrained 174 by stream channel material that would allow full penetration of the temperature probe. Transition 175 from sand / silt to cobble bottom material prevented the extension of temperature maps beyond \sim 176 10 to 25 m from the left bank of the study reach. Areas where groundwater appeared to be closest 177 to the sediment-water interface were selected for deployment of sediment DET and DGT probes.



180 Diffusive gradients in thin-films (DGT) and diffusive equilibrium in thin-films (DET) are 181 methods of measuring fine-scale (cm to mm) solute concentrations in surface waters and sediment 182 pore waters (Davison and Zhang, 2016). They have been used to quantify contaminant 183 concentrations and biogeochemical processes in a variety of environments, including lacustrine 184 (Ma et al., 2020; Zhang et al., 1995), estuarine (Cánovas et al., 2020), marine (Parker et al., 2017), 185 and freshwater sediments (Byrne et al., 2015). Typically, DET devices are used to measure 186 equilibrium surface water and sediment pore water concentrations by allowing solutes to diffuse 187 through a membrane layer (0.45 µm) into an inner hydrogel layer (usually 0.12 cm thick). The 188 time to reach equilibrium depends on the thickness of the DET material diffusion layer (combined 189 hydrogel and membrane layer thickness), the difference between concentrations inside and outside 190 the diffusion layer, and the rate of diffusion. Under the deployment conditions of this study (U diffusion coefficient: 3.11×10^{-6} cm² s⁻¹ for sediment pore waters and 4.44×10^{-6} cm² s⁻¹ for stream 191 192 water based on different pH; Hutchins et al. (2012)), the U concentration in a freshly deployed 193 DET probe is expected to have reached 95% of the surrounding water concentration at 124 min. 194 After deployment, the solute in the hydrogel can be sliced into segments (usually 0.2 to 1 cm) 195 that can be eluted and analyzed for solutes of interest. The major advantage of the technique over 196 other in-situ methods of sediment pore water sampling (e.g. drive points, dialysis peepers) is that

measurements can be made at a higher spatial (vertical) resolution, although the sampling depth is typically restricted to 15 cm by the geometry of the commercially available probe housing (DGT Research Ltd., www.dgtresearch.com). In addition, DET probes typically sample a very small volume of sediment pore water (1 cm depth intervals = 0.1 cm x 1 cm x 1.8 cm = ~ 0.18 cm³) perpendicular to the probe interface. This means there is very little disturbance or averaging of 202 pore water concentrations and chemical gradients, compared to other pore water sampling methods203 (Stockdale et al., 2009).

204 The DGT technique measures the diffusive flux of solute from sediment pore water by 205 introducing a localized sink for the solute in the form of an ion-binding layer that is separated from 206 the sediment pore water by a well-defined material diffusion layer consisting of a filter and 207 hydrogel. Following deployment, a linear diffusion gradient is rapidly ($\leq 60 \text{ min}$) formed across 208 the diffusion layer and solute progressively accumulates in the binding layer (Davison and Zhang, 209 2016). The binding layer can then be analyzed to show spatial differences in the solute fluxes 210 across the probe interface. As the DGT measures the flux of solute into the resin at a given location, 211 it can provide highly localized information about solute mobilization / sequestration processes over 212 the deployment period (Lehto, 2016). Moreover, because the resin layer mostly binds free ions and 213 ions that can be released by labile or partially labile complexes, it can provide unique information 214 about the speciation and bioavailability of solutes in-situ (Amato et al., 2014; Davison and Zhang, 215 2012; Zhang et al., 1995).

216 For groundwater (shallow sediment) deployments, a network of 10 DGT and 10 DET probes 217 were deployed in pairs (within approximately 30 cm of each other) in August 2017 in the sediment 218 on the left bank of the study reach along the Little Wind River (Figure 1). The sediment probes 219 were 15 cm in length and deployed vertically in the sediment. It is worth noting that deployment 220 of DET and DGT sediment probes is typically constrained to sand-dominated sediments that allow 221 easy insertion of the plastic device into the sediment. However, stainless steel holders have been 222 used to successfully deploy DET probes in coarse riverbed environments (Ullah et al., 2012). The 223 probes were retrieved after 48 hours (DET) and 72 hours (DGT) and processed (see section 2.4) 224 within 2 hours of retrieval at the U.S. Geological Survey laboratory in Riverton, Wyoming. The

225 hydrogel layers in the DET and DGT devices were 0.118 cm and 0.078 cm thick, respectively, and 226 used a 0.014 cm thick polyethersulfone membrane (0.45 µm pore size) to separate the hydrogel 227 from the sediment. Numerical modelling of the lateral diffusion of U (using DIFS: (Harper et al., 228 2000)) within the material diffusion layer found that the mean gel concentrations at distances of 1 229 cm and 2 cm intervals were 60% and 13% of the pore water concentrations at the origin after 48 230 h, respectively. For surface water deployments, DET and DGT 'piston' samplers were situated at 231 three locations on the left bank of the Little Wind River: (1) 500 m upstream and (2) 450 m 232 downstream from the cold water anomaly indicated by streambed temperature mapping, and (3) 233 approximately 100 m downstream from the cold water anomaly and suspected groundwater efflux 234 zone (Figure 1). At surface water sampling sites, DET and DGT probes were deployed in triplicate 235 and retrieved and processed as per the shallow sediment deployments. Surface water 'grab' 236 samples were also collected along river transects that overlapped the location of DET and DGT 237 surface water probes. In this instance, samples were collected from the left, center and right side 238 of each transect. Samples were filtered (0.45 µm) on-site and preserved with 1M HNO₃ to await 239 analysis. Further information on the surface water grab sampling procedure are provided in the 240 supplementary information.

The binding layer utilized in DGT devices used a MetsorbTM (TiO₂) sorbent, which has been
shown to be suitable for U measurements in freshwaters (Turner et al., 2014; Turner et al., 2012).
All DET and DGT probes were supplied by DGT Research Ltd. (Lancaster, UK).

244

245 2.4 Laboratory and analytical procedures

In the laboratory, the sediment DGT and DET probes were sliced into 1 cm sections. The hydrogel and binding layers were removed from the DET and DGT probes, respectively, and

248 eluted for 24 hours in 10 mL 1M HNO₃ to await analysis. The solutions (including surface water 249 grab samples) were then analyzed by ICP-MS for target analytes (U and strontium (Sr)). Analysis 250 by ICP-MS utilized rhodium as an internal standard to compensate for analytical drift. Instrument detection limits (IDL = $3.3\sigma / s$) were 0.004 µg L⁻¹ and 0.8 µg L⁻¹ for U and Sr, respectively. High 251 252 precision and accuracy of the control standards $(\pm 4\%)$ and certified reference material (SLRS-6) 253 $(\pm 7\%)$ was achieved. The DGT results are expressed as solute concentrations (C_{DGT}) to help 254 comparison with the DET results. The standard equations used to calculate C_{DGT} are given in the 255 supplementary information.

256

257 2.5 Validation of DGT probes

258 Accumulation of solutes in the DGT device is known to be limited by the buffering capacity of the sediment or water environment. If the flux [ng cm⁻²] demanded by the DGT is in excess of 259 260 solute re-supply, then C_{DGT} may be substantially lower than solute concentrations derived from 261 DET measurements. To test the relation between the flux demanded by the DGT and the actual 262 flux from surface water and sediment pore waters, five sets of DGT 'piston'-style probes (Model: 263 R-SLU, DGT Research Ltd.) with different material diffusion layer thicknesses (filter and 264 hydrogel: 0.054, 0.092, 0.132, 0.170 and 0.208 cm) were inserted into the sediment (10 cm depth) 265 and suspended in the river water at site WR17-6 for 72 hours (Figure 1), each set consisted of three 266 replicate probes. If the flux from surface water and sediment pore waters meets or exceeds the 267 DGT demand over the length of the deployment, a plot of measured mass per unit area versus the 268 material diffusion layer thicknesses should be linear (Zhang et al., 1995).

269

270 2.6 Geochemical speciation modelling

271 Dissolved U(VI) speciation was calculated using PHREEQC (Parkhurst and Appelo, 2013) for 272 sediment pore waters using the major ion chemistry for minipiezometer samples collected at 15 273 cm depth within 1 m of each DET / DGT deployment site and for surface water using averaged 274 major ion chemistry along the study reach (which varied less than 3% among nine surface water 275 grab sample sites) (data available in Naftz et al., 2019). PHREEQC calculations were conducted 276 at 25 °C using the wateq4 database and included aqueous U species stability constants from 277 Guillaumont et al. (2003). Because of their importance on U speciation and solubility, stability 278 constants for ternary (Ca,Mg)-U(VI)-CO₃ complexes were included (Dong and Brooks, 2006; 279 Dong and Brooks, 2008). Thermodynamic data for adjusting many of the U stability constants to 280 the experimental temperature are not available. Oxic conditions were imposed in the PHREEQC 281 speciation calculations because measured dissolved oxygen levels at 30 cm depth were equal to or 282 greater than 0.4 mg L⁻¹ (Naftz et al, 2019). Dissolved oxygen was not measured at shallower 283 depths, with surface water dissolved oxygen. Dissolved uranium is thus assumed to be all in +6 284 oxidation state, (U(VI)). Dissolved Fe and Mn were not included in speciation calculations.

285

3. Results and Discussion

287 *3.1 Transport of uranium from groundwater to surface water*

288 Contour maps of streambed temperatures were constructed for the sampling period in 2017 289 (Figure 2a). The < 25th percentile of streambed temperature was identified to provide a point of 290 reference to denote areas with a higher potential for groundwater discharge through the streambed. 291 The colder areas of streambed sediment were limited to narrow (< 5 m) areas along the left bank 292 of the active channel to the north of the study area (Figure 2a). Streambed temperature values in 293 the southwest corner of the study area were generally > 25th percentile of the streambed 294 temperature values except for one small area (sample point WR17-1) (Figure 2a).

295 Identification of the groundwater efflux (cold water anomalies) from the streambed 296 temperature survey (Figure 2a) guided the selection of sediment pore water DET and DGT 297 sampling sites. Contour maps of DET U concentrations in the sediment pore waters (top 15 cm) 298 along the Little Wind River study reach during August 2017 are presented in Figure 2b, and 299 concentration statistics are summarized in Table 1. Site WR17-1 (upgradient from the groundwater 300 plume) and site WR17-10 (downgradient from below the groundwater plume) are not shown as mean sediment pore water concentrations of U were low (WR17-1, 22 µg L⁻¹; WR17-10, 11 µg L⁻¹ 301 302 ¹), and these sites appeared not to be influenced by contaminated groundwater. The DET probes 303 clearly identified the location and focus of contaminated groundwater in the shallow sediments of 304 the Little Wind River. The shallow portions of the plume appeared to be constrained to an 305 approximately 140 m distance along the channel, its approximate boundaries aligning with sites 306 WR17-9 (upstream) and WR17-8 (downstream). The mean background U concentration (from site WR17-1) was 22 µg L⁻¹. However, the mean U concentration within the groundwater plume zone 307 was 570 μ g L⁻¹ (range = 30 to 1321 μ g L⁻¹), with the peak concentrations centered on site WR17-308 309 6. Direct comparison of these shallow sediment pore water concentrations with other sites is not 310 possible; however, comparison with groundwater samples from Riverton (1 m depth) (Naftz et al., 311 2019) and other UMTRCA sites (> 1 m depth) (Green River, L-Bar, Naturita, Shiprock, and Tuba 312 City, Figure 1) (U.S. Department of Energy Legacy Management, 2020) indicates the DET U concentrations are within the general range reported (13 μ g L⁻¹ to 7100 μ g L⁻¹). 313 314 Between sites WR17-4 and WR17-7, U concentrations decreased towards the surface above

approximately 6 to 7 cm depth (allowing for uncertainty arising from lateral diffusion in the probe),

potentially limiting surface water contamination. Decreases in sediment pore water solutes in near-316 317 surface riverbed sediments has been linked to either physical dilution with infiltrating and lower 318 concentration surface water (Byrne et al., 2014), or sorption or precipitation of mineral phases 319 along a biogeochemical gradient (Fuller and Harvey, 2000). To investigate the process driving 320 attenuation of U in the shallow sediments, we consider Sr concentrations here as an indicator of 321 the degree of groundwater-surface mixing. Strontium is generally considered to behave 322 conservatively in groundwater-surface water systems, and so any decrease in concentration in the 323 sediment pore waters are likely related to mixing with surface waters that would contain lower Sr 324 concentrations (Petelet-Giraud et al., 2018). Figure 2c illustrates a contour map of Sr pore water 325 concentrations from the DET samplers. Unlike U concentrations, elevated Sr pore water 326 concentrations at sites WR17-5, -6 and -7 (range: 4.6 to 7.4 mg L⁻¹) persisted closer to the surface (surface water = $1.1 \text{ mg } \text{L}^{-1}$), with some evidence for decreases at site WR17-7 (to 2.8 mg L^{-1}). 327 328 These slightly different patterns in reactive and conservative solute concentrations indicate both 329 dilution by infiltrating low U concentration surface water (Table 1) and reactive uptake of U in 330 groundwater by sediments during hyporheic mixing may account for the observed decrease in U 331 concentrations. However, the specific geochemical and hydrological processes driving the 332 observed solute concentrations and patterns are not resolved in this study but are currently being 333 investigated.

The effect of the groundwater plume on U concentrations in the Little Wind River is illustrated in Figure 3. Both surface water grab and DET samples showed a longitudinal increase in U concentrations from upstream from the plume to the approximate central location of the plume (Figure 3a and 3b). Mean grab sample U concentrations were lower than the U-DET concentrations and also indicated an increase of U concentrations downstream from the plume area where the 339 DET samples showed a decrease. In this instance, the DET samplers were located closer (within 1 340 m) to the left bank of the Little Wind River than the grab samples and so probably represent higher 341 concentration and poorly mixed plume-derived waters 'hugging' the left bank of the river. As the 342 grab samples were collected from different locations across river transects, the mean of these 343 values incorporates the effect of mixing and is therefore a better representation of longitudinal 344 changes in U concentrations in this relatively wide river channel. Although DET samples were not 345 taken in the same manner as grab samples on this occasion, the DET data still demonstrate a change 346 in surface water U concentrations associated with contaminated groundwater efflux to the river. A 347 longitudinal increase in surface water U concentrations from upstream to downstream from the 348 plume area was also demonstrated in 2016 under lower river flow conditions (see supplementary 349 information and Figure S1) and, as far as the authors are aware, this is the first time that 350 groundwater with elevated U has been shown to increase river water U concentrations at 351 UMTRCA sites. This is despite regular and long-term monitoring of surface waters by DOE and 352 may indicate that the DOE sampling strategy at UMTRCA sites is not representative of locations 353 of contaminated groundwater discharge.

354

355 *3.2 Exploring uranium speciation in sediment pore waters and surface water*

Sediment pore water and surface water C_{DGT} U concentrations were substantially lower than concentrations derived from DET measurements (Table 1); the mean C_{DGT} for U across all ten sample sites was 15 µg L⁻¹ (range: 1 to 122 µg L⁻¹). It is not uncommon for sediment pore water and surface water concentrations derived from DET, drive point samplers, or grab samples to be higher than C_{DGT} (Davison and Zhang, 2012). Typically, there are three possible causes for this. First, re-supply of solute from sediment pore water to the DGT may not be sufficient to meet the demand from the DGT (Lehto, 2016). Second, the binding gel may not be able to remove the target solute from solution rapidly due to competition for binding sites with other solutes (Bennett et al., 2016). Third, the solute of interest may not exist in available forms and are instead partially labile or inert to DGT; this has been used to imply potential constraints to solute bioavailability (i.e. solute that could potentially be absorbed and retained by an organism) (DeGryse and Smolders, 2016). Complexes with large dissolved organic ligands often have small diffusion coefficients and can be less labile than dissolved inorganic complexes (Davison and Zhang, 2012).

369 We can investigate the possibility of solute demand from the DGT exceeding solute supply 370 from the sediment pore waters by calculating $R_{c,72hr}$, the ratio of C_{DGT} from a 72 hour deployment 371 to the bulk pore water concentrations (from DET measurements) (Figure 4) (Lehto, 2016). 372 Numerical modelling carried out with the DGT-induced fluxes in sediments model (Harper et al., 373 1998; Menezes-Blackburn et al., 2019) (assuming sediment porosity of 0.8), indicates that values 374 of R_{c,72hr} greater than 0.80 would indicate a well-buffered system with continuous supply from the 375 solid- and solution phases at a rate almost equal to the flux demanded by the DGT, while values 376 approaching 0.07 would indicate an increasingly diffusion-driven supply. In this scenario, DGT 377 concentrations are expected to approximate DET concentrations. Using the mean C_{DGT} and DET 378 concentrations across the depth profiles (n = 15), it is clear that many $R_{c,72hr}$ values are below even 379 the threshold for diffusive only supply ($R_{c,72hr} = 0.07$) in these sediments (Figure 4). This may be 380 because of spatial and temporal differences in solute supply / concentrations in the sediments or 381 speciation effects on the DGT measurement. Sediments are famously heterogeneous and small-382 scale variability can confound the direct comparison of DET and DGT measurements carried out 383 in different parts of the sediment (Huang et al., 2019). Interpretation of R_{c,72h} values across the 384 depth profile could also be confounded by localized changes in pore water chemistry that could conceivably affect the lability of U to DGT at different depths. Our measurements did not provide
evidence for such changes in these sediments.

387 The R_{c.72h} values in the surface water were also low and these cannot be explained by spatial 388 heterogeneity. Moreover, the supply of solutes to the DGT are not expected to be limited in flowing 389 surface water to the same extent as in sediments. It is important to recognize that C_{DGT} represents 390 a time-integrated mean concentration across the 72 hour measurement period, whereas the DET 391 concentration only reflects conditions within the ~2 hours preceding the end of the deployment, 392 hence temporal differences in U supply from the sediment through variation in hyporheic water 393 flow and exchange may also be important here. Although the data presented here do not allow for 394 complementary analysis of temporal changes in hyporheic flow conditions or solute supply to 395 sediment pore waters and the overlying river water, evidence of temporal (< 24 hour) shifts in 396 sediment pore water U concentrations was found at this study site (Figure S2).

397 Focusing on solute supply to the DGT in the surface water and sediment deployments, we can 398 examine more closely if the U flux demanded by the DGT was greater than solute supply by 399 deploying DGT probes with different material diffusion layer thicknesses. In both surface water 400 and sediment deployments, a linear relationship between U accumulation and the material 401 diffusion layer thickness is evident for probes with layer thicknesses of 0.092 to 0.208 cm (Figure 402 5), indicating sufficient solute supply to satisfy demand from the DGT probes at 10 cm depth at 403 WR17-6. However, in both the sediment and surface water deployments, probes utilizing the 0.054 404 cm layer thickness deviated from linearity. Evidently, for deployments using the thinnest diffusion 405 layer, which demands a higher flux from sediments to the probe, the local supply of solute was 406 exhausted (Zhang et al., 1995). As the DGT field deployments in this study used a diffusion layer 407 thickness of 0.092 cm, we can assume that solute supply to the DGT probe was not limiting

408 accumulation, and spatial and temporal heterogeneity is a possible factor behind the low $R_{c, 72hr}$ 409 values at this location (Figure 4). However, the reduction in the mass accumulated by the probes 410 with the thinnest diffusion layer in the surface water deployments (Figure 5b) indicates that 411 competition for binding sites on the DGT may be important here, as the thinnest diffusion layer 412 would also allow the greatest fluxes of other solute that can outcompete the U for the resin binding 413 sites, thus reducing the total U uptake. It is therefore possible that U species that are inert, partially 414 labile or not bound by DGT resin gel play an important role.

415 To establish the dominant U species in solution, PHREEQC modelling of aqueous U(VI) was 416 conducted. The analysis indicates U speciation was dominated (>98%) by ternary uranyl (Ca or 417 Mg) carbonate complexes in all sediment pore water at 15 cm depth and in surface water, with the 418 uncharged Ca₂UO₂(CO₃)₃ complex accounting for 66-80% of all aqueous U species (Table S2). 419 This is important as uncharged U complexes are unlikely to bind to the DGT resin but will be 420 measured by DET, thus contributing to the low Rc, 72hr values observed. Furthermore, only a small fraction of the total U concentration is present as the dicarbonato U species $(UO_2(CO_3)_2^{2-})$, which 421 has previously been found to best predict U bioavailability to a model invertebrate (Croteau et al., 422 423 2016).

424 Several studies also point to ion competition as a potential explanation for low U accumulation 425 in DGT probes. In laboratory experiments, and in the absence of competing ligands, Ca (< 250 mg 426 L^{-1}) has been shown to aid U uptake on the DGT MetsorbTM gel by forming labile calcium uranyl 427 species (Turner et al., 2012). However, U uptake was reduced at Ca concentrations > 250 mg L^{-1} . 428 Sediment pore water samples (15 cm depth) at the Riverton site had Ca concentrations 331 to 861 429 mg L^{-1} and surface water averaged 72 mg L^{-1} (Naftz et al., 2019). Yet, DGT probes deployed in 430 both environments had low $R_{c, 72hr}$ values. Other potential U complexants such as SO₄ were 431 reported to have no interference effects on U uptake (Turner et al., 2012) however the test range 432 in that study (0.02 to 200 mg L⁻¹) was below the range of SO₄ concentrations in Riverton surface and sediment pore water (15 cm depth) (204 to 6740 mg L⁻¹) (Naftz et al., 2019). Turner et al. 433 (2012) report U uptake was negatively affected by increasing HCO₃ (0.1 to 500 mg L^{-1}) and PO₄ 434 $(0.005 \text{ to } 5 \text{ mg } \text{L}^{-1})$ concentration. Bicarbonate concentrations were elevated in the Riverton 435 sediment pore waters at 15 cm depth (423 to 755 mg L⁻¹) and surface water (179 mg L⁻¹) (Naftz et 436 437 al., 2019). Drozdzak et al. (2016) reported decreased U uptake on the DGT PIWBA resin in the 438 presence of high Ca and SO₄ concentrations. Although these previous studies were not conducted 439 under the same field conditions as the Little Wind River, they do provide evidence that major ion 440 concentrations (Ca, SO₄, HCO₃) in the sediment pore water and surface water were at high enough 441 concentrations to slow U accumulation in the DGT probes due to competition effects.

442 A final factor potentially limiting U accumulation in the DGT resins was the presence of U-443 dissolved organic matter (DOM) complexes. Dissolved organic matter is well known to limit the 444 toxicity of certain metals (Paller et al., 2019). This effect is represented in DGT by the small 445 diffusion co-efficient and partial lability of many DOM-metal complexes (Davison and Zhang, 446 2012). Whilst U-DOM species have been shown to limit U bioavailability to some freshwater 447 invertebrate species (Croteau et al., 2016), the effect of U complexation by DOM in the present study, which ranges from 5 mg C L⁻¹ in surface water to > 30 mg C L⁻¹ in the contaminated 448 449 groundwater at 1 m depth (Naftz et al., 2019), could not be considered in the PHREEQC speciation 450 modelling. This is because the binding constants for U complexation by the DOM at this site are 451 unknown. Therefore, the extent of U complexation by DOM in the Little Wind River groundwater 452 and surface water system could not be examined.

454 3.3 Implications for monitoring U in groundwater-surface water systems

455 A growing number of studies worldwide report elevated concentrations of U in groundwater 456 and drinking water (Banning et al., 2017; Coyte et al., 2018; Nolan and Weber, 2015; Nriagu et 457 al., 2012). With many global regions demonstrating a climate shift towards aridity (Garreaud et 458 al., 2020; Kogan and Guo, 2015), groundwater will become an increasingly more important source 459 of water for humans (Dalin et al., 2017) and have a greater influence on surface water quality 460 (Lansdown et al., 2015). It is therefore critical to develop and test methods to establish: (1) the 461 degree of connectivity between groundwater and surface water systems, and (2) the potential water 462 quality and ecological effects of contaminated groundwater on surface water systems.

463 In this study, streambed temperature surveys combined with DET and DGT groundwater 464 (shallow sediment) and surface water measurements of solute chemistry have been demonstrated 465 as a reliable method of establishing the location of diffuse (U-rich) groundwater efflux to surface 466 water, and the magnitude of the effect on surface water and ground water quality. Sediment pore 467 water contamination was substantial when mean plume U concentrations are compared to 468 background concentrations and drinking water standards. However, a decrease in U pore water 469 concentrations occurred above approximately 6 cm depth, hypothesized to be a combination of 470 shallow groundwater mixing with lower concentration surface water, and precipitation and / or 471 sorption of U solutes to the sediment. These processes could have a substantial bearing on 472 ecosystem health as the hyporheic zone (defined here as the zone of surface water and ground 473 water mixing) has been widely documented as an important habitat and refugium for aquatic 474 organisms (Krause et al., 2011; Stubbington et al., 2011). Furthermore, although the groundwater 475 plume appeared to increase U concentrations in the receiving surface water environment, surface 476 water U concentrations remained below the drinking water standard. Therefore, surface water 477 quality effects from U-rich groundwater plumes may be negligible if the hydrological and 478 geochemical environment in the riverbed sediments promotes attenuation through geochemical 479 reaction, and if the dilution capacity of the surface water system is sufficient to maintain U 480 concentrations below environmental quality standards.

481 The findings of this study indicate aqueous U in the Little Wind River had limited 482 bioavailability. This is inferred from low accumulation of U in the DGT sediment pore water and 483 surface water probes and the dominance of the uncharged Ca₂UO₂(CO₃)₃ aqueous complex 484 determined by speciation modelling. This is consistent with low concentrations of U measured in 485 aquatic invertebrates sampled at this site (Naftz et al., 2019). Considering the dominance of 486 uncharged $Ca_2UO_2(CO_3)_3$ at the Riverton site, the high concentrations of solutes known to compete 487 with U for DGT binding sites, and the inferred low U bioavailability from DGT investigations, 488 analysis of groundwater and surface water chemistry at other UMTRCA sites allows us to offer 489 some initial insights on the potential bioavailability of U at these sites. Figure 6 illustrates a 490 comparison of groundwater and surface water chemistry at seven UMTRCA sites across the 491 western U.S.A (data obtained from GEMS database; U.S. Department of Energy Legacy 492 Management, 2020). Similar to Riverton, water chemistry is broadly characterized as calcium, 493 bicarbonate and sulfate-type, with high total dissolved solids (240 12000 mg L⁻¹) and high 494 concentrations of ions (Ca, SO₄, Na, HCO₃) known to compete with U for biological uptake sites 495 in DGT probes and model organisms. Although we have not performed PHREEQC speciation 496 analyses of these waters, or deployed DGT probes, it is reasonable at this stage to hypothesize that 497 U bioavailability might be limited at these sites through complexation and / or competition effects. 498 However, further investigations at the highlighted UMTRCA sites would be needed in order to 499 verify this hypothesis.

501 4. Conclusions

502 Growing concerns over U contamination of groundwater, combined with increasing relevance 503 of groundwater resources for drinking water and riverine ecosystems, have highlighted the need 504 to understand the environmental risk of U in groundwater-surface water systems. To this end, this 505 study has presented a methodological framework to establish the transport of U from groundwater 506 to surface water and its potential ecological effects. We applied this methodology in the Little 507 Wind River where U-rich groundwater is interacting with surface water downgradient from the 508 former Riverton U mill and tailings disposal site. This site is typical of many former mill sites in 509 the western U.S.A. now under the management of the DOE. Streambed temperature mapping was 510 initially used to identify a broad zone of groundwater discharge to the river. Diffusive equilibrium 511 in thin-film (DET) probes were then used to map vertical and longitudinal patterns in sediment 512 pore water U and Sr concentrations. This further constrained the zone of groundwater discharge, 513 and demonstrated decreases in U concentrations in the shallow sediment pore waters through 514 groundwater-surface water interaction processes. An increase in river U concentrations was 515 observed in the area of groundwater discharge; however, concentrations did not exceed 516 environmental guidelines. This is the first time that contaminated groundwater at UMTRCA sites 517 has been shown to affect surface waters and emphasizes the importance of a carefully designed 518 and targeted sampling strategy to establish the effect of groundwater plume discharge on surface 519 waters. Low accumulation of U in diffusive gradients in thin film (DGT) samplers, and the 520 dominance of the uncharged Ca₂UO₂(CO₃)₃ complex, indicates limited bioavailability of this 521 element in this river system. Further, we hypothesize low U bioavailability at other former mill 522 sites in the western U.S.A. due to similar groundwater and surface water chemistry. However, the

523 extent of U complexation by DOM was not explored in this study and further investigation of the 524 role of DOM in aqueous U speciation and DGT measurements is warranted. Application of the 525 methodological framework used in this study at other DOE and worldwide sites with legacy 526 groundwater issues may be highly beneficial to establish the extent of surface water and ecosystem 527 contamination in groundwater-surface water systems.

- 528
- 529 CRediT authorship contribution statement

530 Patrick Byrne: Conceptualization, Methodology, Investigation, Formal analysis, 531 Visualization, Writing – original draft, Writing – review and editing. Christopher C. Fuller: 532 Conceptualization, Methodology, Formal analysis - geochemical modelling, Writing - review and 533 David L. Naftz: Conceptualization, Methodology, Writing - review and editing, editing. 534 Supervision. Robert L. Runkel: Conceptualization, Methodology, Formal analysis, Writing – 535 review and editing. Niklas J. Lehto: Methodology, Formal analysis - DIFS modelling, Writing -536 review and editing; William L. Dam: Writing – review and editing.

537

538 **Declaration of competing interest**

539 The authors declare that they have no known competing financial interests or personal 540 relationships that could have appeared to influence the work reported in this paper.

541

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550 Figure and table captions

Figure 1. Study area on the Little Wind River, Riverton, Wyoming, U.S.A., showing all DET and DGT sediment and surface water sampling sites. Top left inset shows location of selected UMTRCA sites in the USA. Bottom right inset shows the location of the former Riverton Processing Site and the approximate boundary and flow direction of the U-rich groundwater plume.

Figure 2. (a) Streambed temperature profile along the left bank of the Little Wind River from
August 2017. Sediment pore water uranium (b) and strontium (c) concentrations obtained from
DET measurements along the left bank of the Little Wind River.

Figure 3. Comparison of surface water uranium concentrations from upstream to downstream from the location of groundwater entry to the Little Wind River, showing (a) grab samples, (b) DET measurements, and (c) DGT measurements. Error bars in Figure 3a are the standard deviation of three samples collected at the left side, center, and right side of a transect along the river. Error bars in Figures 3b and 3c are the standard deviation of triplicate measurements from the left bank of the river only. Riv-U/S = sample site upstream from the plume. Riv-Mid = sample site at approximately the location of the plume. Riv-D/S = sample site downstream from the plume.

566 Figure 4. Accumulation of uranium in sediment (WR17) and surface water (Riv) DGT probes 567 represented as $R_{c, 72hr}$, the ratio of C_{DGT} from a 72 hour deployment to the bulk pore water solute

concentrations from DET measurements. $R_{c, 72hr} > 0.8$ = sustained case; $0.07 < R_{c, 72hr} > 0.8$ = partially sustained case; $R_{c, 72hr} < 0.07$ = diffusive case. Error bars are the standard deviation of 15 measurements for sediment samplers and 3 measurements for surface water samplers. Vertical solid lines represent the approximate upstream and downstream boundaries of the groundwater plume entry to the Little Wind River. Samples are shown from upstream (left) to downstream (right).

574 Figure 5. Measured mass per unit area of uranium plotted against the reciprocal of the material 575 diffusion layer thickness (Δg) in the DGT probes deployed in (a) shallow sediments (10 cm depth) 576 and (b) surface waters for 72 hrs. Error bars are the standard deviation of triplicate measurements.

577 Figure 6. Piper plot showing the major ion chemistry of ground and surface water at selected 578 UMTRCA sites, including the former Riverton Processing site. Data represent the most recent 579 sample available from a single measurement site on the GEMS database [U.S. Department of 580 Energy Legacy Management, 2020]. GW = groundwater sample. SW = surface water sample.

Table 1. Summary of uranium concentrations in µg L⁻¹ from DET and DGT deployments and surface water grab samples in the Little Wind River. Data for surface water (SW) grab samples are concentrations at the start and end of surface water DET and DGT deployments. Data for sediment pore water (PW) samples are mean, minimum, and maximum concentrations.

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586 **References**

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Figure 1. Study area on the Little Wind River, Riverton, Wyoming, U.S.A., showing all DET and DGT sediment and surface water

- sampling sites. Top left inset shows location of selected UMTRCA sites in the USA. Bottom right inset shows the location of the former
- 776 Riverton Processing Site and the approximate boundary and flow direction of the U-rich groundwater plume.





Figure 2. (a) Streambed temperature profile along the left bank of the Little Wind River from August 2017. Sediment pore water uranium







Figure 3. Comparison of surface water uranium concentrations from upstream to downstream from the location of groundwater entry to the Little Wind River, showing (a) grab samples, (b) DET measurements, and (c) DGT measurements. Error bars in Figure 3a are the standard deviation of three samples taken at the left side, centre, and right side of a transect along the river. Error bars in Figures 3b and 3c are the standard deviation of triplicate measurements from the left bank of the river only. Riv-U/S = sample site upstream of the plume. Riv-Mid = sample site at approximately the location of the plume. Riv-D/S = sample site downstream of the plume.

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Figure 5. Measured mass per unit area of uranium plotted against the reciprocal of the material diffusion layer thickness (Δg) in the DGT probes deployed in (a) shallow sediments (10 cm depth) and (b) surface waters for 72 hrs. Error bars are the standard deviation of triplicate measurements.





801 Figure 6. Piper plot showing the major ion chemistry of ground and surface water at selected UMTRCA sites, including the former

- 802 Riverton Processing site. Data represent the most recent sample available from a single measurement site on the GEMS database [U.S.
- 803 Department of Energy Legacy Management, 2020]. GW = groundwater sample. SW = surface water sample.

805	Table 1. Summary of uranium concentrations in μ g L ⁻¹ from DET and DGT deployments and surface water grab samples in the Little
806	Wind River. Data for surface water (SW) grab samples are concentrations at the start and end of surface water DET and DGT
807	deployments. Data for sediment pore water (PW) samples are mean, minimum, and maximum concentrations.

	Riv-U/S	Riv-Mid	Riv-D/S
Grab-SW $(n = 2)$	5.1 / 5.5	6.8 / 6.5	6.3 / 6.4
DET-SW $(n = 3)$	6.7 (6.3 – 6.9)	9.6 (7.7 – 12.6)	7.5 (7.4 – 7.8)
DGT-SW $(n = 3)$	0.2 (0.13 – 0.24)	0.18 (0.16 - 0.21)	0.32 (0.32 – 0.33)

	WR17-1	WR17-9	WR17-2	WR17-3	WR17-4
DET-PW (n = 15)	21.5 (19.4 - 23.5)	4.8 (3.9 – 6.1)	179 (67 – 227)	351 (282 - 441)	794 (718 – 834)
DGT-PW (n = 15)	2.8 (1.9 - 4.4)	0.7 (0.3 – 1.6)	6.9 (2.3 – 14.5)	8.3 (6 - 13.5)	28.6 (15 – 47.4)

	WR17-5	WR17-6	WR17-7	WR17-8	WR17-10
DET-PW (n = 15)	903 (610 - 1049)	1010 (487 – 1321)	675 (449 - 861)	77 (31 – 148)	11.1 (8.9 – 14.3)
DGT-PW (n = 15)	19.6 (14.1 – 29.6)	56 (20.8 - 121.7)	21 (13.8 - 35.4)	11.5 (4.4 – 29)	1.8 (1.5 - 2.3)