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1 **Isotopic Evidence for Mercury Photoreduction and Retention on Particles in Surface**
2 **Waters of Central California, USA**

3
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13

14 **Abstract:**

15

16 Cache Creek (Coast Range, California) and the Yuba River (Sierra Nevada Foothills,

17 California) are two river systems affected by extensive mercury (Hg) contamination due to

18 legacy sources of Hg related to mining. Stable Hg isotope techniques have proven useful for

19 elucidating the complex cycling of Hg within aquatic ecosystems, and we applied these

20 techniques to improve understanding of Hg and methylmercury (MeHg) transformations in these

21 watersheds. Total mercury (THg) concentrations and Hg stable isotope ratios were measured in

22 filtered surface waters and suspended particulate matter collected from 14 sites within the Cache

23 Cr. and Yuba R. watersheds. Filtered surface waters from both watersheds exhibited values of

24 $\Delta^{199}\text{Hg}$ (0.37‰ to 0.71‰), consistently elevated above those observed in sediments ($\Delta^{199}\text{Hg}$

25 average = 0.07‰). Associated suspended particulates from these surface water samples

26 displayed a much greater range of values for $\Delta^{199}\text{Hg}$ (-0.61‰ to 0.70‰), although suspended

27 particulates from the Yuba R. exhibited mostly negative $\Delta^{199}\text{Hg}$ values (-0.61‰ to 0.10‰). The

28 relationship between $\Delta^{199}\text{Hg}$ and $\Delta^{201}\text{Hg}$ in the filtered surface waters and associated suspended

29 particulates was calculated using a bivariate York regression, yielding a slope of 1.57 ± 0.49

30 ($\pm 2\text{SE}$) for the Yuba R. and 1.40 ± 0.27 ($\pm 2\text{SE}$) for Cache Cr., both within error of the previously

31 reported experimentally-derived slopes for MeHg- and inorganic Hg(II)- photoreduction. This

32 provides isotopic evidence that Hg photoreduction is occurring within these surface waters to a
33 significant degree, and suspended particulate phases are retaining the reduced product of Hg
34 photoreduction, particularly within the Yuba R. The isotopic compositions of filtered surface
35 waters are consistent with the isotopic signatures recorded in biota at low trophic positions
36 within these watersheds, suggesting that the reservoir of Hg incorporated within the biota of
37 these systems is similar to the filter-passing Hg fraction in surface waters.

38

39 **Keywords:** mercury; stable mercury isotopes; mercury photoreduction; contaminated fluvial
40 ecosystem

41

42 **1. Introduction**

43 Historic mining activities in the watersheds of numerous Central Valley, California rivers
44 have resulted in an enduring legacy of widespread mercury (Hg) contamination throughout the
45 region. In the Sierra Nevada foothills, hydraulic gold mining activities in the nineteenth century
46 mobilized immense amounts of Hg-contaminated sediment into the downstream portions of
47 rivers like the Yuba River, and the effects of these activities continue to impact river function
48 and ecosystem health both locally and far downstream (Domalgalski, 2001; Marvin-DiPasquale
49 et al., 2003; Domalgalske et al., 2004; Alpers et al., 2005; James et al., 2009; Bouse et al., 2010;
50 Ghoshal et al., 2010; Fleck et al., 2011; Marvin-DiPasquale et al., 2011 Springborn et al., 2011;
51 Donovan et al., 2013; Singer et al., 2013; Higson and Singer, 2015; Singer et al., 2016;
52 Nakamura et al., 2018). In the California Coast Range, Hg-ores have been mined since the mid-
53 nineteenth century to produce metallic Hg, creating substantial contamination related to release
54 of Hg from the retorting process and mine waste tailings in the rivers that drain these areas, such

55 as Cache Cr. (Domalgalski et al., 2004 USGS; Marvin-DiPasquale et al., 2009; Suchanek et al.,
56 2010). Coast Range watersheds also experience continuing inputs of Hg from natural processes
57 related to hydrothermal activity and leaching from Hg-containing outcrops (Smith et al., 2008
58 and references therein; Suchanek et al., 2010). The continuing impacts of this widespread
59 contamination on biota in these systems, as evidenced by elevated methylmercury
60 concentrations, has been well documented for both aquatic and terrestrial organisms (May et al.,
61 2000 USGS; Alpers et al., 2004 USGS; Hothem et al., 2007; Hothem et al., 2008; Hothem et al.,
62 2010; Fleck et al., 2011 USGS) and extends downstream into the San Francisco Estuary (Eagles-
63 Smith et al., 2009; Greenfield & Jahn, 2010; Donovan et al., 2013). There is, however, a
64 significant knowledge gap within the field of Hg biogeochemistry as to how elevated sources of
65 inorganic mercury (IHg), such as the legacy mining sources affecting the Yuba R. and Cache Cr.
66 watersheds, correspond to and affect methylation of Hg and subsequent bioaccumulation within
67 organisms (Domagalski et al., 2004).

68 More specifically, there are still open questions about the transformation of aqueous Hg
69 species into the bioavailable pool of Hg within freshwater fluvial ecosystems (Ward et al., 2010).
70 Stable Hg isotope studies have emerged within the last decade that demonstrate the ability of this
71 method to elucidate the complex transformations Hg undergoes during environmental cycling
72 (Yin et al., 2010; Blum et al., 2014; Blum & Johnson, 2017). Specifically, Hg isotopes have been
73 used within California river systems to study the relationship between sediment-associated Hg
74 and concentrations of Hg within invertebrates and fish (Donovan et al., 2016a; Donovan et al.,
75 2016b). These studies identified distinct MeHg and IHg isotope endmembers within the study
76 watersheds, but have been unable to identify specific processes that could link these two
77 endmembers. Analysis of the specific isotopic shifts between IHg and MeHg led investigators to

78 suggest that two processes, photoreduction of MeHg and biotic fractionation (a balance of
79 methylation and demethylation processes), were necessary to explain the Hg isotopic
80 composition of biota. Experimental studies of photochemical reduction of both IHg and MeHg
81 have demonstrated that these processes impart characteristic odd mass independent fractionation
82 (MIF) signatures (Bergquist and Blum, 2007; Zheng and Hintelmann, 2009; Chandan et al.,
83 2015; Rose et al., 2015), but the limited number of field studies conducted in freshwater aquatic
84 ecosystems to date have focused on the isotopic composition of receptor organisms (e.g.
85 macroinvertebrates, fish) rather than abiotic Hg pools (Tsui et al., 2013; Kwon et al., 2015;
86 Lepak et al., 2018).

87 Using stable Hg isotope analysis, this study builds upon previous research by
88 investigating the transformation of Hg within surface waters, particularly photoreduction
89 processes, to understand the role these species play in the isotopic shifts associated with
90 transformation of IHg to bioavailable forms. California surface waters exhibit significant odd
91 MIF anomalies, providing isotopic evidence that photoreduction processes are actively affecting
92 Hg cycling in these fluvial systems. California surface waters also display isotopic compositions
93 that are consistent with the isotopic compositions of low trophic position biota within these
94 watersheds, providing evidence that the reservoir of bioavailable MeHg is also present in the
95 surface water pool.

96 **2. Materials and Methods**

97 **2.1 Sample Collection and Processing**

98 Filtered stream water and suspended sediment samples were collected at fifteen locations
99 along the Yuba R. (including the Feather River downstream of the confluence with the Yuba R.),
100 Cache Cr. (including Bear Creek and Sulfur Creek in the headwater regions of Cache Cr.), and

101 the Yolo Bypass Wildlife Area (YBWA) during June 2015 (06/02/15 to 06/09/15). Sampling
102 locations within each watershed were chosen to align with sites at which samples had been
103 previously collected and analyzed for Hg isotopic analysis by Donovan et al. (2016a) and
104 Donovan et al. (2016b) (Figure S6). Water samples ranging from 1 L to 15 L in volume were
105 collected, filtered, and preserved in the field, using trace-metal clean sampling methods
106 following a modification of EPA Method 1669. All filtered water field blanks (n=8) had THg
107 concentrations that were below method detection limits of 0.2 ng/L. At each site, 1L of water
108 was collected into a HDPE bottle and used to determine the total suspended solids (TSS) of
109 surface water. TSS values were used in the calculation of distribution coefficients ($\log K_d$) using
110 THg values of the associated filtered surface water and suspended material following the method
111 of Hurley et al. (1998).

112 **2.2 Sample Preparation for Isotope Analysis and THg Concentrations**

113 Hg in suspended material was separated for THg concentration and Hg stable isotope
114 measurement by offline combustion, as described in detail elsewhere (Biswas et al., 2008;
115 Demers et al., 2013). Hg in filtered surface water was separated for Hg stable isotope
116 measurement by adsorption onto an anion exchange resin (AG-1X4, Biorad, 200-400 mesh)
117 using a method modified from the procedure described in Štok et al. (2014) and detailed in the
118 unabridged methods [SI S1]. Prior to anion exchange, THg concentrations of each surface water
119 sample were determined by analysis of small aliquots of UV-treated samples via CV-AFS
120 (following EPA method 1631). Recovery of filtered surface water samples during the anion
121 exchange resin column process averaged $91.6\% \pm 3.1\%$ (range: 86.2% to 95.5%, n=7). Three
122 samples with suspected high DOC contents had poor recoveries (YBWA PW#2, 4.1%; Rumsey,
123 Cache Cr, 58.8%; Bear Cr. @HCC, 47.9%); the mass independent isotopic composition of

124 samples is unchanged by incomplete yields, but the mass dependent isotopic composition may
125 have been affected. Therefore, although reported in this manuscript, the mass dependent isotopic
126 composition of these samples is suspect, and the $\Delta^{xxx}\text{Hg}$ values of these low-recovery samples
127 are displayed in manuscript figures and tables with an asterisk. While not an exhaustive
128 investigation, these low sample yields of Hg suggest that the anion exchange resin column
129 method may not be the ideal pre-concentration protocol for low Hg concentration waters with
130 high DOC contents.

131 Trapping solutions of both combustion and column resin-exchange samples were
132 partially reduced with a 30% solution of $\text{NH}_2\text{OH}\cdot\text{HCl}$ using an amount equal to 2% of the total
133 sample by weight (w/w); then a small aliquot was taken and measured for THg by CV-AFS or
134 cold vapor atomic absorption spectroscopy [CV-AAS; MA-2000 Nippon Instruments].
135 Combustion trap contents and surface water eluents were then purged into a secondary 1%
136 KMnO_4 trapping solution to remove potential matrix components and to adjust Hg
137 concentrations prior to isotopic analysis (Sherman and Blum, 2013; Blum and Johnson, 2017).
138 Transfer recoveries into secondary traps for concentration and matrix- matching for all sample
139 types averaged $96.8\% \pm 5.8\%$ (range: 73.3% to 106.6%, $n=42$, only 2 transfers $<85\%$ recovery).
140 None of these transfers are likely to have significantly fractionated the processed samples (Blum
141 and Johnson, 2017).

142 Procedural blanks were processed in parallel with samples for THg concentration and Hg
143 isotopic composition. The trap contents of combusted field filter blanks contained 104 ± 18 pg of
144 Hg ($n=8$, $\pm 1\text{SD}$), which is not significantly different from procedural combustion blanks
145 (105 ± 74 pg of Hg, $n=10$, $\pm 1\text{SD}$). Column resin-exchange procedural blanks yielded between 119
146 and 191 pg of Hg ($n=4$, $\text{mean} \pm 1\text{SD} = 149 \pm 30$ pg), representing no more than 5.6% of Hg in

147 sample eluent solutions.

148 2.3 Hg Isotope Analysis

149 The Hg isotopic composition of the secondary trapping solutions were measured by cold
150 vapor multi-collector inductively coupled plasma mass spectrometry (CV-MC-ICP-MS) at either
151 the University of Michigan (Nu Instruments) or the University of Toronto (Thermo Fisher
152 Neptune Plus). Mercury stable isotope compositions are reported throughout this paper in permil
153 (‰) using delta notation ($\delta^{\text{xxx}}\text{Hg}$) relative to NIST SRM 3133 (Eq. 1), with mass dependent
154 fractionation based on the $^{202}\text{Hg}/^{198}\text{Hg}$ ratio ($\delta^{202}\text{Hg}$) (Blum and Bergquist, 2007). Mass
155 independent fractionation is reported as the deviation from the theoretically predicted $\delta^{\text{xxx}}\text{Hg}$
156 values based on the kinetic mass fractionation law and is reported with capital delta notation
157 ($\Delta^{\text{xxx}}\text{Hg}$) according to Eq. 2. In this study MIF is represented with $\Delta^{199}\text{Hg}$, $\Delta^{200}\text{Hg}$, $\Delta^{201}\text{Hg}$, and
158 $\Delta^{204}\text{Hg}$, using $\beta = 0.252$, $\beta = 0.502$, $\beta = 0.752$, and $\beta = 1.493$, respectively (Blum and Bergquist,
159 2007).

160 Equation 1: $\delta^{\text{xxx}}\text{Hg} (\text{‰}) = \left(\left[\frac{(^{\text{xxx}}\text{Hg}/^{198}\text{Hg})_{\text{Sample}}}{(^{\text{xxx}}\text{Hg}/^{198}\text{Hg})_{\text{NIST3133}}} \right] - 1 \right) \times 1000$

161 Equation 2: $\Delta^{\text{xxx}}\text{Hg} (\text{‰}) = \delta^{\text{xxx}}\text{Hg} - (\delta^{202}\text{Hg} \times \beta)$

162 Certified reference materials and standards were processed and analyzed along with the
163 California water samples. UM-Almáden was used as the process reference material for the
164 column resin exchange procedures. Column anion exchange resin recoveries for UM-Almáden
165 averaged $97.3\% \pm 3.7\%$ (range: 93.3% to 101.7%, n=4). NIST 2711 “Montana Soil,” was chosen
166 as a suitable process reference material for offline combustions due to its high THg conc.
167 ($6.25\mu\text{g/g}$) and similar matrix. Combustion recoveries of NIST 2711 averaged $102.3\% \pm 5.8\%$
168 (range: 92.6% to 108.9%, average THg conc = $6391 \pm 365 \mu\text{g/g}$, n=8). The average isotopic
169 composition of NIST 2711 ($\delta^{202}\text{Hg} = -0.25 \pm 0.10\text{‰}$, $\Delta^{199}\text{Hg} = -0.27 \pm 0.03\text{‰}$) was consistent

170 with previously reported values (Estrade et al., 2011; Jiskra et al., 2015; Yin et al., 2016; Blum
171 and Johnson, 2017; Washburn et al., 2017; Washburn et al., 2018a; Washburn et al., 2018b).
172 UM-Almáden was measured during each analytical session on the CV-MC-ICP-MS to quantify
173 within-session performance. The isotopic data from these process reference materials and
174 process standards is summarized in Table S1. The analytical uncertainty of Hg isotopic
175 measurements (2SD) is presented regardless of sample preparation method used, because the
176 error associated with the 2SD of UM-Almáden (averaged across analytical sessions) was greater
177 for all measured Hg isotope values than that associated with the sample matrix specific process
178 reference material (2SE). Hence, the analytical uncertainty of Hg for samples in this study is
179 presented as $\delta^{202}\text{Hg} \pm 0.12\text{‰}$, $\Delta^{199}\text{Hg} \pm 0.03\text{‰}$, $\Delta^{201}\text{Hg} \pm 0.08\text{‰}$, $\Delta^{200}\text{Hg} \pm 0.05\text{‰}$ and $\Delta^{204}\text{Hg} \pm$
180 0.20‰ . [Table S1].

181 **3. Results and Discussion**

182 **3.1 Isotopic Composition of Hg in California Surface Waters**

183 Filtered surface waters collected from the Yuba R. and Feather R. displayed low THg
184 concentrations (0.48 to 0.86 ng/L), and relatively small variation in isotopic composition ($\delta^{202}\text{Hg}$
185 = -0.82‰ to -0.50‰, avg. = $-0.69 \pm 0.13 \text{‰}$ [1SD]; $\Delta^{199}\text{Hg}$ = 0.37‰ to 0.62‰, avg. = 0.49 ± 0.14
186 ‰ [1SD]) [Table 1; Figure 1]. The associated suspended particulates displayed a much greater
187 range in both THg concentrations (166.1 to 665.2 ng/g) and isotopic composition ($\delta^{202}\text{Hg}$ = -
188 1.31‰ to 0.25‰, avg. = $-0.68 \pm 0.54\text{‰}$ [1SD]; $\Delta^{199}\text{Hg}$ = -0.61‰ to 0.10‰, avg. = $-0.21 \pm$
189 0.24‰ [1SD]) [Table 1; Figure 1]. Both filtered surface waters [Unpaired t-Test with equal
190 variance, $n_1=4$, $n_2=12$, $T=11.40$, $p < 0.001$] and suspended particulates [Unpaired t-Test with
191 equal variance, $n_1=6$, $n_2=12$, $T=3.731$, $p=0.003$] have significantly different average $\Delta^{199}\text{Hg}$
192 values than Yuba R. riverbank and terrace sediments ($\Delta^{199}\text{Hg} = 0.04 \pm 0.03 \text{‰}$ [1SD]) (Donovan

193 et al., 2016a).

194 Filtered surface waters collected from Cache Cr. and Bear Cr. (which flows into Cache
195 Cr.) displayed relatively low THg concentrations (1.03 to 11.78 ng/L), and relatively limited
196 variation in isotopic composition ($\delta^{202}\text{Hg} = -0.84\text{‰}$ to -0.73‰ ; $\Delta^{199}\text{Hg} = 0.58\text{‰}$ to 0.64‰). This
197 small range likely reflects the low number of data points analyzed for Hg isotope composition
198 ($n=2$) [Table 1; Figure 2]. Similar to the Yuba R., the associated suspended particulates
199 displayed a much greater range in both THg concentration (114.6 to 2788 ng/g) and isotopic
200 composition ($\delta^{202}\text{Hg} = -1.78\text{‰}$ to -0.91‰ , avg. = $-1.23 \pm 0.30\text{‰}$ [1SD]; $\Delta^{199}\text{Hg} = 0.05\text{‰}$ to
201 0.53‰ , avg. = $0.19 \pm 0.17\text{‰}$ [1SD]) although this greater range may be influenced by the greater
202 number of suspended particulate samples analyzed for Hg isotope composition ($n=7$) [Table 1;
203 Figure 2]. Unlike the Yuba R., Cache Cr. suspended particulates did not have a significantly
204 different average $\Delta^{199}\text{Hg}$ value compared to riverbed and terrace sediments ($\Delta^{199}\text{Hg} = 0.10 \pm$
205 0.06‰ [1SD]) [Unpaired t-Test with equal variance, $n_1=7$, $n_2=14$, $T=1.804$, $p=0.087$] (Donovan
206 et al., 2016b). Filtered surface waters from Cache Cr. displayed $\Delta^{199}\text{Hg}$ values that were
207 significantly elevated compared to sediments [Unpaired t-Test with equal variance, $n_1=2$, $n_2=14$,
208 $T=267$, $p << 0.001$]. Filtered surface water and suspended particulates collected from Sulfur
209 Creek displayed the greatest THg concentration (179.1 ng/L), the most negative $\delta^{202}\text{Hg}$ values ($-$
210 2.47‰ and -2.89‰ , respectively) and the most positive $\Delta^{199}\text{Hg}$ values (0.71‰ and 0.70‰ ,
211 respectively) of any surface waters analyzed in this study [Figure 2, Table 1].

212 **3.2 Isotopic Evidence for Mercury Photoreduction in Surface Waters**

213 **3.2.1 $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ Ratio in Surface Waters**

214 Filtered surface waters from both watersheds exhibited consistently elevated $\Delta^{199}\text{Hg}$
215 values (0.37‰ to 0.71‰) and $\Delta^{201}\text{Hg}$ values (0.21‰ to 0.51‰) (Figure 4; Table 1). The

216 associated suspended particulates from these surface waters displayed a much greater range of
217 $\Delta^{199}\text{Hg}$ values (-0.61‰ to 0.70‰) and $\Delta^{201}\text{Hg}$ values (-0.38‰ to 0.50‰), but a number of
218 particulate samples had negative MIF values, suggesting suspended particulates may be affected
219 by a different set of fractionation processes, or are retaining a different phase of Hg than filtered
220 surface waters (Figure 3, Figure 4, Figure S1, Figure S5, Table 1).

221 Previous experimental studies have established that the $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ ratio of samples
222 can be diagnostic of the occurrence of specific fractionation processes, including photochemical
223 reduction reactions (Bergquist & Blum, 2007; Zheng & Hintelmann, 2009; Rose et al., 2015;
224 Chandan et al., 2015). If a singular fractionation mechanism was responsible for the odd mass-
225 independent isotopic compositions of the CA surface waters, one would expect that they would
226 exhibit characteristic $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ ratios in good agreement with one of the previously
227 determined experimental slopes for MIF inducing mechanisms. Regression statistics for the
228 relationship between $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ in the filtered surface water and associated suspended
229 particulates of each river were calculated using the Model 2 bivariate York regressions
230 embedded in *Isoplot* (v.3.00, Ludwig) [Figure 4] and found to be 1.57 ± 0.49 ($\pm 2\text{SE}$) for the Yuba
231 River and 1.40 ± 0.27 ($\pm 2\text{SE}$) for Cache Creek. Applying the same regression model to the entire
232 surface waters dataset presented in this manuscript yields a slope that is statistically
233 indistinguishable from the slopes calculated for the rivers individually: 1.52 ± 0.24 ($\pm 2\text{SE}$) (Figure
234 4). The $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ slopes for each river overlap with a number of previously reported,
235 experimentally-derived slopes. These include MeHg photo-reduction with a slope of 1.36 ± 0.08
236 ($\pm 2\text{SE}$) (Bergquist and Blum, 2007), photoreduction of Hg(II) at environmentally relevant
237 Hg/DOC ratios with slopes ranging from 1.19 ± 0.02 to 1.31 ± 0.14 ($\pm 2\text{SE}$) (Zheng & Hintelmann,
238 2009), and an experimentally-derived slope of 1.59 ± 0.05 ($\pm 2\text{SE}$) for nuclear volume effects

239 (NVE)-related fractionation associated with processes such as abiotic reduction by DOM (Zheng
240 and Hintelmann, 2010). Given that each river as well as the overall surface water dataset is
241 potentially consistent with a number of MIF-inducing mechanisms, it is possible that odd-MIF
242 isotopic composition of surface waters in these rivers may not be controlled by a single process.

243 If just considering filtered surface water samples, the calculated $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ York
244 regression slope of $1.13 \pm 0.32 (\pm 2\text{SE})$ is significantly lower than the overall dataset slope. Filtered
245 surface water samples exhibit MIF isotopic compositions that are consistent with both MeHg
246 photoreduction and photoreduction of Hg(II) at environmentally relevant Hg/DOC ratios (Zheng
247 & Hintelmann, 2009; Bergquist and Blum, 2007; Chandan et al., 2015). Photoreduction of both
248 MeHg and Hg(II) can produce the relatively large magnitude MIF anomalies observed in the
249 filtered surface water samples ($\Delta^{199}\text{Hg}$ values up to 0.71‰) with much smaller MDF shifts.

250 Although not directly measured in this study, MeHg concentrations in filtered surface waters of
251 the Yuba R. and Cache Cr. watersheds have previously been demonstrated to be low, with MeHg
252 concentrations typically $\ll 1\text{ng/L}$ and representing $<10\%$ of THg of these surface waters
253 (Domalgalski et al., 2004 USGS; Hunerlach et al., 2004; Stumpner et al., 2018). Despite the low
254 MeHg concentrations, the odd-MIF signatures of Hg in the surface waters of these watersheds
255 suggests that the Hg remaining in these surface waters is partially influenced by MeHg
256 photoreduction processes. A combination of these two photoreduction processes, acting on IHg
257 and MeHg reservoirs with slightly different starting MIF isotopic compositions, could produce
258 the observed $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ ratio in the surface waters we have analyzed (Figure S5).

259 The large degree of uncertainty in the calculated regression $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ slope for
260 filtered surface waters could be driven by differences in the starting isotopic composition of pre-
261 photodegraded Hg (either MeHg or inorganic Hg(II)) between the two watersheds, as well as

262 spatial variations within each watershed. Previous studies have suggested that the differences in
263 the isotopic composition of pre-photodegraded MeHg could be related to contrasting net biotic
264 fractionation (a balance between biotic methylation and demethylation processes) between sites,
265 or the large range in IHg source isotopic composition observed in these two watersheds
266 (Donovan et al., 2016a; Donovan et al., 2016b). Neither filtered surface waters nor suspended
267 particulates exhibit significant longitudinal trends in $\Delta^{199}\text{Hg}$ values along the sampling transects
268 in the Yuba R. or Cache Cr. (Fig. S3). In both watersheds there is sufficient variation in
269 suspended particulate $\Delta^{199}\text{Hg}$ values between adjacent sampling locations such that the transport
270 of particulates downstream could influence the overall isotopic composition in downstream
271 filtered surface water samples if sufficient exchange of Hg occurred between these phases.

272 The $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ slope of filtered surface waters [$1.13 \pm 0.32 (\pm 2\text{SE})$] does not agree
273 with the experimentally derived slope for NVE-related MIF [$1.59 \pm 0.05 (\pm 2\text{SE})$], but including
274 the particulate samples in the $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ slope linear regression shifts the regression slopes
275 for each river to within error of NVE effects. As the filtered surface water and suspended
276 particulate Hg phases are likely undergoing parallel fractionation processes, it is possible that the
277 MIF anomalies observed in filtered surface waters are partially created by NVE-related
278 processes. However, NVE has only been observed to produce small odd-MIF signatures of \leq
279 $+0.40\text{‰}$ in simulated natural systems (Zheng & Hintelmann, 2010), which is less than the
280 magnitude observed in the surface water samples ($\Delta^{199}\text{Hg}$ up to 0.71‰). Importantly, NVE-
281 related shifts in $\Delta^{199}\text{Hg}$ would also impart much larger shifts in $\delta^{202}\text{Hg}$ values for the filtered
282 surface waters (shifts up to $\sim 4.00\text{‰}$), and such shifts are not observed in any of the filtered
283 surface water samples – the greatest $\delta^{202}\text{Hg}$ shift observed for any filtered surface water sample
284 from the river sediment average is -0.44‰ (Simpson Bridge site). Hence, it is unlikely that NVE-

285 related processes are responsible for the observed $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ ratio in surface waters or are
286 driving the large magnitude odd-MIF anomalies.

287 Suspended particulate samples from both watersheds are in general agreement with the
288 $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ slope values predicted for the individual river sample sets, indicating that the
289 odd-MIF signatures of Hg within these samples is partially produced by the same photoreduction
290 fractionation process. Suspended particulates collected from sites within the Yuba R. watershed
291 exhibit mostly negative odd-MIF signatures ($\Delta^{199}\text{Hg} = -0.61\text{‰}$ to 0.10‰ , mean = $-0.21 \pm 0.24\text{‰}$
292 [$\pm 1\text{SD}$]), while associated suspended particulates collected from the Cache Cr. watershed exhibit
293 consistently positive odd-MIF signatures ($\Delta^{199}\text{Hg} = 0.05\text{‰}$ to 0.53‰ , mean = $0.19 \pm 0.17\text{‰}$
294 [$\pm 1\text{SD}$]). The different signs of the odd-MIF anomalies of suspended particulates for Yuba R.
295 versus Cache Cr. suggests that there are differences in the photochemical processing of Hg
296 within the two watersheds.

297 **3.2.2 Retention of Hg Photodegradation Products on Suspended Particulates**

298 In the Yuba R., suspended particulate phases appear to be retaining the reduced product
299 of MeHg and Hg(II) photoreduction, as this reduced product would have significantly more
300 negative $\Delta^{199}\text{Hg}$ values than the starting MeHg/Hg(II) pool (Zheng & Hintelmann, 2009).
301 Retention of photoreduced Hg(0) could be occurring due to oxidation and subsequent aqueous
302 sorption either on or within the particles. Direct oxidation of Hg(0) by reduced DOM under
303 anoxic conditions has been observed experimentally (Gu et al., 2012), and such reduction could
304 be occurring within anoxic microenvironments of particulate organic matter aggregates
305 (Böckelmann et al., 2000; Ortiz et al., 2015). A recent study by Zheng et al., (2018),
306 demonstrated experimentally that the Hg(II) fraction produced during oxidation by organic
307 matter is shifted to slightly more negative $\Delta^{199}\text{Hg}$ values ($\sim -0.20\text{‰}$) than the starting Hg(0)

308 reservoir due to NVE-related effects. Hence, the oxidation of photoreduced Hg(0) would
309 enhance the negative MIF signatures within the suspended particulate phase by further shifting
310 the isotopic composition of the retained Hg fraction to more negative odd MIF values (Figure
311 S5).

312 Other processes could also be contributing to the negative $\Delta^{199}\text{Hg}$ values observed in
313 Yuba R. suspended particulates. Soils in the watershed of the Yuba R. have been shown to have
314 slightly negative $\Delta^{199}\text{Hg}$ values ($\Delta^{199}\text{Hg} = -0.1$ to -0.06%) (Zheng et al., 2016). If a significant
315 portion of the Hg pool found in suspended particulates is sourced from watershed inputs of Hg
316 associated with allochthonous organic matter, this could result in slightly negative $\Delta^{199}\text{Hg}$ values
317 (Jiskra et al., 2017), but probably not of the magnitude observed in the Yuba R. samples. Such
318 mixing would also result in shifts to lower $\delta^{202}\text{Hg}$ values that are not observed in suspended
319 particulates, making this explanation unlikely. Abiotic reduction of Hg(II) by organic matter
320 associated with the suspended particulates would result in the Hg(II) retained on the particulates
321 to be shifted to more negative $\Delta^{199}\text{Hg}$ values, with a shift on the order of -0.20% to -0.60%
322 (Zheng and Hintelmann, 2010). Abiotic reduction of Hg(II) could be contributing to the overall
323 isotopic signature within suspended particulates, although it is unlikely that is the major process
324 contributing to the negative odd MIF signatures of Yuba R. suspended particulates because
325 relatively large magnitude MIF shifts would also induce a significantly larger magnitude positive
326 MDF shift (up to $\sim +5.50\%$) that was not observed in suspended particulate samples. Another
327 process that could result in negative $\Delta^{199}\text{Hg}$ values is oxidation of elemental Hg present in the
328 sediment fraction as a result of past mining activity, although a large pool of Hg(0) would need
329 to be present in the surface waters to create an observable shift in odd MIF signatures.

330 The consistently positive $\Delta^{199}\text{Hg}$ values in Cache Cr. suspended particulates contrasts

331 with observations from the Yuba R. One explanation for this difference is that riverbed and
332 terrace sediments in Cache Cr. (sediments represent the largest source of IHg to surface waters in
333 the channel) have significantly more positive $\Delta^{199}\text{Hg}$ values on average ($0.10\pm 0.06\%$ [$\pm 1\text{SD}$])
334 than those observed in the Yuba R. ($0.04\pm 0.03\%$ [$\pm 1\text{SD}$]) [Unpaired t-Test with equal variance,
335 $n_1=12$, $n_2=14$, $T=2.87$, $p<0.01$] (Donovan et al., 2016a; Donovan et al., 2016b). Suspended
336 particulates may be retaining photoreduced Hg with negative $\Delta^{199}\text{Hg}$ values, but this signal is
337 diluted after mixing with IHg that has positive $\Delta^{199}\text{Hg}$ values. Another possible explanation for
338 the more positive $\Delta^{199}\text{Hg}$ values in Cache Cr. suspended particulates is that retention of
339 photoreduced Hg is more limited in Cache Cr, perhaps due to differing composition of particles
340 within the system (e.g. differing mineral compositions with lower affinity for Hg sorption, lower
341 density of thiol-like binding sites within organic matter substrates, more limited aggregation of
342 particles). If the pool of Hg being photoreduced in Cache Cr. is predominantly bound to S-
343 bearing ligands within organic matter complexes, experimental work suggests that the produced
344 Hg(0) would actually have positively shifted $\Delta^{199}\text{Hg}$ values, which if retained would result in a
345 $\Delta^{199}\text{Hg}$ value in the suspended particulates that was positively shifted compared to the associated
346 sediments (Zheng and Hintelmann, 2010). Zheng and Hintelmann (2010) showed, however, that
347 Hg bound to bulk dissolved organic matter did not exhibit the MIF signatures associated with
348 lower molecular weight S-bearing organic complexes, suggesting that MIF related to
349 photoreduction of S-bearing organic compounds may be a relatively limited process in the Cache
350 Cr. watershed.

351 Four suspended particulate samples from Cache Cr. have $\Delta^{199}\text{Hg}$ values that are more
352 positive than the riverbed and terrace sediment average. For these samples, largely located near
353 headwater Hg sources associated with hydrothermal and mining activity, the MIF signatures of

354 suspended particulates likely represents the predominance of the source contributions.
355 Additionally, mixing with IHg sourced from the hydrothermal and mining activities in the
356 headwaters may have resulted in a more positive $\Delta^{199}\text{Hg}$ value in pre-photodegraded MeHg and
357 Hg(II), particularly for samples collected from Bear Cr.

358 **3.2.3 Extent of Mercury Photoreduction Processes**

359 The degree to which MeHg or inorganic Hg(II) has been photodegraded within a given
360 sample can be calculated from the isotopic composition of that sample by assuming a starting
361 isotopic composition of the MeHg/Hg(II) pool subject to prior photochemical degradation. The
362 average $\Delta^{199}\text{Hg}$ value within sediment from each watershed (Yuba R. $\Delta^{199}\text{Hg} = 0.04 \pm 0.03\%$;
363 Cache Cr. $\Delta^{199}\text{Hg} = 0.10 \pm 0.06\%$; compiled from Donovan et al., 2016a and Donovan et al.,
364 2016b) can be used as the starting isotopic composition of both pre-photodegraded MeHg and
365 Hg(II), because the net biotic processes that produce MeHg do not induce MIF (Kritee et al.,
366 2009; Rodriguez-Gonzalez et al., 2009). By assuming that all MIF is related to MeHg
367 photodegradation and subtracting the sediment average $\Delta^{199}\text{Hg}$ values from those observed in the
368 filtered surface waters, the total magnitude of $\Delta^{199}\text{Hg}$ shift related to MeHg photodegradation can
369 be estimated. Calculated $\Delta^{199}\text{Hg}$ shifts range from $\Delta^{199}\text{Hg} = 0.33$ to 0.61% for the Yuba R. and
370 Cache Cr. watersheds. Since DOC concentrations in both watersheds are typically between 1-5
371 mg/L (Chow et al., 2007; Domalgalski et al., 2004), experimental data for isotopic fractionation
372 of MeHg during photodegradation in the presence of 1mg/L DOC, was used to estimate the
373 proportion of MeHg in the filtered surface waters subject to photodegradation prior to
374 incorporation in the water column (Bergquist & Blum, 2007). These calculations show that
375 between 10% and 17% of the total mass of Hg in the filtered surface water would have
376 undergone MeHg photochemical degradation to yield the observed $\Delta^{199}\text{Hg}$ shifts. Similarly,

377 experimental data for isotopic fractionation of Hg(II) during photoreduction in the presence of
378 1.2 mg/L DOC was used to estimate the proportion of Hg(II) remaining in filtered surface waters
379 that had been subjected to photodegradation prior to incorporation in the water column (Zheng &
380 Hintelmann, 2009). This calculation yields an estimate that the Hg mass measured in the filtered
381 surface waters represents 74% to 85% of the Hg mass that would have been present prior to
382 photoreduction processes.

383 As %MeHg (percent of THg present as MeHg) values for filtered surface waters of these
384 watersheds are typically <10%, insufficient MeHg would be available to undergo
385 photodegradation to produce the magnitude of MIF signatures observed, supporting the
386 hypothesis that Hg(II) photoreduction processes are contributing to the observed MIF signatures.
387 Additionally, the calculated proportion of MeHg in 2015 surface waters is lower than the
388 calculated degree of photodegradation for the estimated bioavailable MeHg endmember in both
389 the Yuba R. (2013: 24% photodegraded; 2014: 35% photodegraded) and Cache Cr. (2013: 17%
390 photodegraded; 2014: 31% photodegraded). This evidence suggests that the bioavailable MeHg
391 pool present in surface waters is not exclusively contributing to MIF shifts (Donovan et al.,
392 2016a; Donovan et al., 2016b). The lesser extent of photodegradation calculated for 2015 filtered
393 surface water may be related to a number of factors, including differences in water quality
394 parameters (e.g. increased turbidity, increased water depth) that would result in net decreases of
395 photoreduction, or mixing with colloidal-IHg with near-zero $\Delta^{199}\text{Hg}$ values from sediment or
396 suspended particulate sources, resulting in a measured $\Delta^{199}\text{Hg}$ value with a smaller magnitude,
397 thereby biasing our calculated extent of photodegradation to artificially low values.

398 The amount of photoreduced Hg(0) that would need to be retained in the particulate
399 phase of the Yuba R. can be estimated using a simple mixing model calculation. Using the

400 average $\Delta^{199}\text{Hg}$ value for Yuba R. sediments (0.04‰) and the average $\Delta^{199}\text{Hg}$ value for
401 photoreduced $\text{Hg}(0)$ (-3.88‰) from the $\text{Hg}(\text{II})$ -DOM photoreduction experiments conducted by
402 Zheng and Hintelmann (2009), the calculation predicts that to obtain the average $\Delta^{199}\text{Hg}$ value
403 observed for Yuba R. suspended particulates (-0.21‰) only 7% of the total Hg present in the
404 particulates needs to be retained $\text{Hg}(0)$. With only 7% of the Hg present in the suspended
405 particulates (~10 to 20 ng/L) this is a much greater mass of Hg than is found in any of the filter-
406 passing Hg pools in the Yuba R., so this calculation may not accurately account for a factors
407 such as photoreduction of suspended particulate material as well as a potentially greater
408 magnitude of MIF shifts associated with MeHg -DOM photoreduction rather than exclusively
409 $\text{Hg}(\text{II})$ -DOM photoreduction.

410 **3.3 Sources of Hg to the Surface Waters of the Yuba R. and Cache Cr. Watersheds**

411 **3.3.1 Variations in the Isotopic Composition of Filtered Surface Waters and Suspended** 412 **Particulates**

413 Both MeHg and $\text{Hg}(\text{II})$ photoreduction processes impart $\delta^{202}\text{Hg}$ shifts, so surface water
414 Hg pools should have distinct isotopic compositions from the source Hg pools as the result of
415 these processes. For samples collected from the Yuba R., filtered surface water samples and
416 suspended particulates do have significantly different average $\Delta^{199}\text{Hg}$ values [Unpaired t-Test
417 with equal variance, $n_1=6$, $n_2=4$, $T=5.333$, $p=0.001$], but do not have significantly different
418 $\delta^{202}\text{Hg}$ values [Unpaired t-Test with equal variance, $n_1=6$, $n_2=4$, $T=0.035$, $p=0.936$], despite the
419 much greater variation in isotopic composition observed in suspended particulates.

420 For the Yuba R. watershed surface waters, no clear trends are observed between THg
421 concentration and either $\delta^{202}\text{Hg}$ values or $\Delta^{199}\text{Hg}$ values. The partition coefficients ($\log(k_D)$)
422 remain essentially constant throughout the sampling reach (5.62 to 5.72), suggesting that Hg

423 partitioning between the particulate phases and filter-passing phases is similar throughout the
424 study reach, and that Hg inputs from sediment resuspension of bed/bank sediment are similar at
425 each sampling location. Additionally, there are no clear trends between isotopic composition and
426 sampling distance downstream. Furthermore, all samples were collected under the same
427 hydrologic flow regime, so there does not appear to be a temporal bias that could affect Hg
428 partitioning.

429 The lack of correlation between either sample location or THg concentration and the Hg
430 isotopic composition within surface waters suggests that the Hg isotopic composition within the
431 Yuba R. surface waters is not controlled by isotopic end-member mixing or site-specific
432 fractionation processes at any given sampling location. Neither the filtered surface waters nor
433 suspended particulate fractions have average $\delta^{202}\text{Hg}$ values that are significantly different from
434 the average $\delta^{202}\text{Hg}$ values observed in river bank and terrace sediments in the Yuba R. ($\delta^{202}\text{Hg} =$
435 $-0.38 \pm 0.42 \text{ ‰ [1SD]}$) (Donovan et al., 2016a). The similar ranges and average $\delta^{202}\text{Hg}$ values of
436 surface water fractions and sediments support the interpretation that river bank and terrace
437 sediments are the main source of Hg to the surface water in the Yuba R. However, the
438 differences in average $\Delta^{199}\text{Hg}$ values in these three sample types imply that photodegradation
439 fractionation processes result in odd-MIF anomalies in the surface waters, but not in the bank
440 and terrace sediments.

441 In Cache Cr., filtered surface water samples and suspended particulates have average
442 $\delta^{202}\text{Hg}$ values that differ by 0.44‰, but the Hg isotopic composition of these two Hg pools is not
443 significantly different [Unpaired t-Test with equal variance, $n_1=7$, $n_2=2$, $T=2.012$, $p=0.084$].
444 While not statistically significant, this pattern of positive $\delta^{202}\text{Hg}$ shifts is consistent with
445 fractionation due to photoreduction processes. Similar to the Yuba R., the two sample types do

446 have significantly different average $\Delta^{199}\text{Hg}$ values [Unpaired t-Test with equal variance, $n_1=7$,
447 $n_2=2$, $T=3.321$, $p=0.013$]. No clear trends between THg concentration and either $\delta^{202}\text{Hg}$ values or
448 $\Delta^{199}\text{Hg}$ values are observed in the Cache Cr. surface waters. THg concentrations for both filtered
449 surface water and suspended particulates increase with distance downstream, but this increase is
450 not reflected in clear trends in Hg isotopic composition [Table 1]. Hg partition coefficients are
451 generally lower in Cache Cr. and they increase with distance downstream ($\log(k_D) = 5.18$ to
452 5.44), which may be indicative of increasing inputs of sediment-sourced Hg in the particulate
453 phase. The riverbed and terrace sediments in Cache Cr. had a more positive average $\delta^{202}\text{Hg}$ value
454 ($\delta^{202}\text{Hg} = -0.84 \pm 0.49\text{‰}$ [1SD]) than was observed in suspended particulates, but this difference
455 was not significant [Unpaired t-Test with equal variance, $n_1=7$, $n_2=14$, $T=1.933$, $p=0.068$]
456 (Donovan et al. 2016b). Overall, there are no clear patterns in $\delta^{202}\text{Hg}$ values for Hg in surface
457 water pools in either the Yuba R. or Cache Cr. In other words, there is no good support for the
458 hypothesis that photoreduction processes exhibit a control on the isotopic compositions of
459 surface water Hg in these systems, while $\Delta^{199}\text{Hg}$ values for these same fractions do provide
460 evidence for this hypothesis. Significant heterogeneity in sediment isotopic composition between
461 sampling sites is likely contributing to this apparent decoupling between MDF and MIF
462 signatures in the surface water Hg pools.

463 **3.3.2 Influence of Sediment Hg on Surface Water Hg Isotopic Composition**

464 The relationship between filtered surface water, associated suspended particulates, and
465 bed or bank sediment samples collected at the same location (subsequently referred to as
466 “paired” samples) is shown in Figure 3. No consistent patterns are observable in the relationship
467 between $\delta^{202}\text{Hg}$ values for paired sample types from the same sampling location. Comparing
468 sediments to suspended particulates, both positive and negative $\delta^{202}\text{Hg}$ shifts ($+0.75\text{‰}$ to -

469 0.82‰) are observed for paired samples. Filtered surface waters also show both positive and
470 negative $\delta^{202}\text{Hg}$ shifts (+0.30‰ to -0.56‰) compared to paired sediments.

471 The apparent decoupling between the isotopic composition of Hg in paired sediments and
472 surface water compartments could be due to a number of factors. Surface water Hg fractions
473 could represent the influence of Hg incorporated at upstream locations and transported
474 downstream. Both Cache Cr. and Yuba R. sediments displayed relatively large ranges in $\delta^{202}\text{Hg}$
475 values (Cache Cr.: -1.69‰ to -0.11‰; Yuba R.: -0.95‰ to 0.71‰). Transport of sediment-
476 bound Hg with a differing isotopic composition from upstream locations could account for the
477 $\delta^{202}\text{Hg}$ shifts observed between sediments at a specific location and suspended particulates. Our
478 sampling campaign was not conducted with sufficient spatial resolution to capture reach-scale
479 heterogeneity in Hg isotopic composition, and there may be significantly more variation than
480 previously documented.

481 As has been demonstrated in the limited number of studies that have investigated the
482 isotopic composition of suspended particulate phase Hg in surface waters (Foucher et al., 2013;
483 Washburn et al., 2017; Jiskra et al., 2017; Washburn et al., 2018b; Demers et al., 2018; Baptista-
484 Salazar et al., 2018), there are a number of processes, such as sorption to mineral phases or
485 aqueous photochemical reduction of Hg, that can produce significant $\delta^{202}\text{Hg}$ shifts in this Hg
486 pool relative to a Hg source (Jiskra et al., 2012; Bergquist and Blum, 2007). A number of
487 sampling site-specific factors, such as turbidity, solar insolation, and biotic activity, could all
488 alter the degree to which certain fractionation processes were occurring at any site. As both
489 positive and negative isotopic shifts were observed from sediment to suspended particulates, no
490 single fractionation processes can be invoked to explain the variation in surface water data.

491 $\Delta^{199}\text{Hg}$ values of co-located “paired” samples of filtered surface water, suspended

492 particulates and stream sediment exhibit similar behavior at each sample site. All suspended
493 particulate samples had lower $\Delta^{199}\text{Hg}$ values compared to sediments, except for the Rumsey site,
494 where the suspended particulates are within uncertainty of the sediment value (suspended
495 particulates $\Delta^{199}\text{Hg} = 0.12\text{‰}$, sediment $\Delta^{199}\text{Hg} = 0.09\text{‰}$). All filtered surface water samples
496 showed more positive $\Delta^{199}\text{Hg}$ values than co-located sediment samples. Unlike the lack of
497 pattern in $\delta^{202}\text{Hg}$ values for paired samples, the consistent pattern observed for $\Delta^{199}\text{Hg}$ values is
498 suggestive of a single fractionation process controlling the odd-MIF composition of the surface
499 water samples, namely Hg photodegradation as discussed in depth in Section 3.2.

500 **3.3.3 Influence of Hydrothermal Activity on the Isotopic Composition of Hg in Sulfur** 501 **Creek**

502 The isotopic composition of these samples is likely impacted by the hydrothermal activity
503 and hot springs (Wilbur Springs) in the headwaters of this stream. The isotopic composition of
504 the surface waters we measured in Sulfur Creek are distinctly different than those reported in
505 Smith et al., 2008 for hydrothermal spring precipitates collected from Wilbur Springs ($\delta^{202}\text{Hg} = -$
506 0.95‰ , $\Delta^{201}\text{Hg} = 0.11\text{‰}$), but they are within the range of $\delta^{202}\text{Hg}$ values for other hydrothermal
507 spring precipitates from the same region of the Coast Range hydrothermal complex ($\delta^{202}\text{Hg} = -$
508 0.95‰ to -3.42‰ , $\Delta^{201}\text{Hg} = 0.11\text{‰}$ to 0.32‰). The relatively more negative $\delta^{202}\text{Hg}$ values
509 observed in Sulfur Creek compared to the Wilbur Springs hydrothermal precipitates could be the
510 result of a temporally variable hydrothermal Hg source to the hot springs, or the result of
511 fractionation related to the precipitation of Hg from the hydrothermal fluid as suggested by
512 Smith et al. (2008). The large magnitude odd-MIF signatures observed in the surface waters of
513 Sulfur Cr. are likely a result of both the elevated above regional background odd MIF signatures
514 of the hydrothermally-derived Hg due to hydrothermal processing, and subsequent

515 photoreduction processes occurring after the hydrothermally-derived Hg reaches the surface
516 environment (discussed in Section 3.2).

517 To determine the influence of Sulfur Creek on the Hg isotopic composition of Bear
518 Creek, surface water samples were obtained both above and below the confluence of Sulfur Cr.
519 with Bear Cr. Below the confluence (Bear Cr. at Holston Chimney Canyon), the filtered surface
520 water and suspended particulate fractions both exhibited elevated THg concentrations compared
521 to measurements taken above the confluence, providing evidence for an influx of Hg from Sulfur
522 Cr. into Bear Cr. The $\delta^{202}\text{Hg}$ value for suspended particulates collected at Holston Chimney
523 Canyon ($\delta^{202}\text{Hg} = -1.03\text{‰}$) was plotted versus 1/THg to evaluate mixing relationships between
524 particulates from Sulfur Cr. ($\delta^{202}\text{Hg} = -2.89\text{‰}$) and Bear Cr. above the confluence with Sulfur
525 Cr. ($\delta^{202}\text{Hg} = -1.39\text{‰}$). The samples do not exhibit a significant a linear trend [linear least
526 squares regression, slope = 2389.7 ± 1925.2 , $p = 0.37$], as particulates from Holston Chimney
527 Canyon have a much less negative $\delta^{202}\text{Hg}$ value that cannot be explained due to inputs of Hg
528 from Sulfur Cr. The lack of a linear relationship between the 1/THg concentration data and the
529 isotope data is likely due to the occurrence of fractionation of Hg originating from Sulfur Cr.
530 once it has entered Bear Cr. Although the $\Delta^{199}\text{Hg}$ value observed in particulates at Holston
531 Chimney Canyon ($\Delta^{199}\text{Hg} = 0.53\text{‰}$) is intermediate between Sulfur Cr. ($\Delta^{199}\text{Hg} = 0.70\text{‰}$) and
532 Bear Cr. above the confluence ($\Delta^{199}\text{Hg} = 0.20\text{‰}$), this value is likely affected by fractionation
533 during photoreduction and other MIF-inducing processes, and does not directly provide evidence
534 for source mixing in Bear Cr.

535 **3.3.4 $\Delta^{204}\text{Hg}$ and $\Delta^{200}\text{Hg}$ Signatures of Surface Waters**

536 None of the filtered surface water samples in the Yuba R. or Cache Cr. has even-MIF
537 signatures of significant magnitude (Figure S1). Of the suspended particulate samples, only 5

538 samples have $\Delta^{204}\text{Hg}$ values that are distinguishable from 0.00‰ based on analytical uncertainty
539 ($\pm 0.20\%$), and only two of these samples (Simpson Bridge site on the Yuba R., $\Delta^{204}\text{Hg} = -$
540 0.29% , $\Delta^{200}\text{Hg} = -0.01\%$; Bear Cr. above Sulfur Cr., $\Delta^{204}\text{Hg} = -0.40\%$, $\Delta^{200}\text{Hg} = 0.04\%$ [Table
541 1]) exhibit negative $\Delta^{204}\text{Hg}$ values that would be consistent with Hg derived from un-impacted
542 North American atmospheric precipitation, but these suspended particulates do not exhibit
543 $\Delta^{200}\text{Hg}$ values that are different from 0.00‰ (Gratz et al., 2010; Chen et al., 2012; Demers et al.,
544 2013; Donovan et al., 2013; Cai and Chen, 2015). Two suspended particulate samples (Dantoni
545 and Parks Bar sites on the Yuba R.) have even MIF signatures consistent with observations of
546 un-impacted atmospheric total gaseous Hg (positive $\Delta^{204}\text{Hg}$ values, negative $\Delta^{200}\text{Hg}$ values)
547 (Gratz et al., 2010; Sherman et al., 2010; Demers et al., 2013; Demers et al., 2015; Fu et al.,
548 2016; Yu et al., 2016). These two samples were run at the lowest (0.35 ng/g) and next to lowest
549 (0.5 ng/g) run solution concentrations, and UM-Almadén run concurrently at these
550 concentrations yielded much greater analytical uncertainty (0.35 ng/g run solution:
551 $\Delta^{204}\text{Hg} \pm 0.56\%$; $\Delta^{200}\text{Hg} \pm 0.28\%$, $n=5$), suggesting that the anomalously large magnitude even-
552 MIF signatures observed in these two samples are unlikely to represent natural processes or
553 sources affecting the suspended particulate load at these two locations. Based on the lack of
554 significant even-MIF signatures in the surface water samples, atmospherically-derived Hg is
555 unlikely to be a significant contributor to the total Hg load in surface waters of the studied
556 watersheds. The even-MIF signatures of these surface water samples provide additional evidence
557 that the predominant Hg source within these systems is related to mining-derived activities rather
558 than atmospheric deposition, in agreement with a previous study that estimated the contributions
559 of atmospheric Hg deposition to these watersheds (Domagalski et al., 2016).

560 **3.4 Comparison of Surface Water Hg Isotopic Composition to Biota within the Yuba R.**

561 **and Cache Cr. Watersheds**

562 Utilizing Hg stable isotopes to identify the transformations that lead to a bioavailable
563 pool of Hg within surface water compartments of fluvial systems was a motivation for
564 conducting this work. For both the Yuba R. and Cache Cr. systems, filtered surface water Hg
565 pools have very similar $\delta^{202}\text{Hg}$, $\Delta^{201}\text{Hg}$, and $\Delta^{199}\text{Hg}$ values as the lower trophic position
566 invertebrates reported in Donovan et al., 2016a and Donovan et al., 2016b (Fig. 1, Fig. 2). The
567 concordant isotopic signatures in low trophic position invertebrates and filtered surface waters is
568 suggestive that the Hg reservoirs in both of these sample types have undergone similar
569 environmental processing, imparting similar degrees of isotopic fractionation. It is important to
570 note that for low trophic position biota, a significant portion of the Hg present within their tissues
571 is inorganic Hg (Hg(II) complexes), and that this inorganic Hg is thought to not bioaccumulate to
572 as significant a degree as MeHg. Previous studies have suggested that the MIF signatures of Hg
573 incorporated by biota are unlikely to be fractionated by internal processes (Perrot et al., 2012;
574 Kwon et al., 2013). The lack of MIF fractionation within biota suggests that both Hg(II) and
575 MeHg affected by photoreduction within surface water prior to incorporation within biota would
576 impart elevated odd-MIF signatures to those biota.

577 For both watersheds, the vast majority of biota have $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ ratio values that fall
578 within uncertainty of the slope predicted for surface water samples collected in this study
579 (1.52 ± 0.24) (Fig. S1), evidence that the Hg(II) and MeHg within these organisms is being
580 affected by the same photoreduction processes suggested to be producing the MIF within the
581 surface water Hg pools. In other words, filter-passing MeHg in the water column could be to a
582 significant source of MeHg to food webs in these rivers systems. Furthermore, when the
583 $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ slopes are evaluated for specific sampling sites at which biota, sediment, and

584 surface water samples have all been collected, the site-specific slopes are all within error of the
585 overall $\Delta^{199}\text{Hg}/\Delta^{201}\text{Hg}$ slopes for surface waters in each river (Figure S4). Although the
586 sediment, biota, and surface water samples were collected during different years (they were,
587 however, collected during the same season), it appears that the processes controlling MIF
588 fractionation within these two river systems are relatively constant temporally and spatially
589 within the rivers.

590 In the Yuba R., the range in filtered surface water isotopic composition ($\delta^{202}\text{Hg} = -0.82\text{‰}$
591 to -0.50‰ , $\Delta^{199}\text{Hg} = 0.37\text{‰}$ to 0.62‰) is similar to a subset of invertebrate organisms with
592 between 37%-143% %MeHg ($\delta^{202}\text{Hg} = -0.76\text{‰}$ to -0.47‰ , $\Delta^{199}\text{Hg} = 0.27\text{‰}$ to 0.62‰), and
593 much more positive $\Delta^{199}\text{Hg}$ values than filamentous algae samples with 2-17% %MeHg ($\delta^{202}\text{Hg}$
594 = -0.82‰ to -0.61‰ , $\Delta^{199}\text{Hg} = 0.06\text{‰}$ to 0.16‰) (Donovan et al., 2016a). Similarly, in Cache
595 Cr. the range in filtered surface water isotopic composition ($\delta^{202}\text{Hg} = -0.84\text{‰}$ to -0.73‰ , $\Delta^{199}\text{Hg}$
596 = 0.58‰ to 0.64‰ , excluding the Sulfur Cr. FSW samples) is within the range exhibited by a
597 subset of invertebrate organisms with between 38%-115% %MeHg ($\delta^{202}\text{Hg} = -1.23\text{‰}$ to -0.69‰ ,
598 $\Delta^{199}\text{Hg} = 0.51\text{‰}$ to 0.69‰) (Donovan et al., 2016b). Interestingly, filamentous algae samples
599 from Cache Cr. exhibited a much greater range in $\Delta^{199}\text{Hg}$ values ($\Delta^{199}\text{Hg} = 0.37\text{‰}$ to 1.00‰ ,
600 $\delta^{202}\text{Hg} = -1.15\text{‰}$ to -0.34‰) and %MeHg (20%-60%), which Donovan et al (2016a) attributed
601 to multiple, isotopically distinct pools of IHg and MeHg in the Cache Cr. system. The large
602 range in filtered surface water isotopic composition observed within the headwaters of Cache
603 Cr., including Sulfur Cr., is consistent with the interpretation that there could be a number of
604 isotopically distinct IHg pools that could undergo methylation and incorporation into low trophic
605 position biota. Such an interpretation would necessitate that MeHg photoreduction was occurring
606 predominantly while MeHg was associated with small particles or dissolved in surface waters

607 prior to incorporation into the food chain. At present, there is insufficient supporting data to
608 confidently make such a claim, and future studies are needed to understand this complex portion
609 of the Hg cycle. Future studies should focus on the relationship between low trophic position
610 biota and filtered surface water Hg isotopic composition, particularly as they fluctuate
611 temporally and with hydrologic conditions.

612 In both Cache Cr. and the Yuba R., negative $\delta^{202}\text{Hg}$ shifts were observed between the
613 estimated IHg source and the estimated pre-photodegraded MeHg source (Donovan et al., 2016a;
614 Donovan et al., 2016b). One hypothesis suggested for this negative $\delta^{202}\text{Hg}$ offset was that *in situ*
615 methylation of Hg within sediments, followed by MeHg advection into the water column and
616 transport downstream by flowing water, would produce a local bioavailable MeHg pool that had
617 undergone minimal biotic MeHg degradation, thus exhibiting more negative net biotic
618 fractionation shift (Donovan et al., 2016a; Donovan et al., 2016b). The isotopic compositions of
619 Hg within the surface waters of Cache Cr. and Yuba R. support this hypothesis, as evidenced by
620 the overlapping isotopic composition with biota. The reservoir of MeHg that is bioavailable and
621 being bioaccumulated within the biota of these fluvial systems is isotopically similar to the Hg
622 reservoir within surface waters. The temporal differences in the isotopic composition of the
623 estimated MeHg end-member observed in the previous studies of these river systems, combined
624 with the lack of temporally concurrent biota data to accompany the 2015 surface water samples
625 prevents us from unequivocally concluding that surface waters are the source of MeHg to the
626 aquatic biota in these river systems. However, the concordant isotopic compositions of Hg in
627 filtered surface waters and low trophic position biota is highly suggestive that the MeHg and
628 Hg(II) in the two compartments is from the same source, and accumulation of surface water
629 MeHg by biota is consistent with the isotopic data.

630 **3.5 Future Implications and Conclusions**

631 We applied stable Hg isotope measurement techniques to surface water samples from two
632 California river systems affected by extensive legacy mining-related Hg contamination in order
633 to better elucidate the biogeochemical cycling of Hg on the path to bioavailability. Isotopic
634 evidence, particularly odd-MIF signatures, indicate that Hg photoreduction processes are
635 occurring within Yuba R. and Cache Cr. surface waters to a significant degree. Within the Yuba
636 R., the isotopic composition of suspended particulate phases suggests retention of the reduced
637 product of MeHg and Hg(II) photoreduction may be occurring. Importantly, the similar isotopic
638 compositions of filtered surface waters and low trophic position biota within these watersheds
639 indicates that the reservoir of Hg within the biota of these California rivers is similar to the filter-
640 passing fraction of Hg in surface waters. Future studies should utilize the Hg preconcentration
641 methods such as the one employed here to analyze other low Hg concentration freshwaters to
642 expand the understanding of the biogeochemical cycle of Hg in aquatic ecosystems that had
643 previously been analytically difficult or unfeasible.

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662 **Supporting Information**

663 The Supporting Information includes an unabridged methods section, and six figures, and
664 one table.

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Figures & Tables

Figure 1

Plot of $\delta^{202}\text{Hg}$ (‰) vs $\Delta^{199}\text{Hg}$ (‰) values for samples from the Yuba River watershed, including one site on the Feather River. Filtered surface waters (red circles) and suspended particulates (blue squares) were collected in June 2016, while values for biota (green triangles) and riverbank and terrace sediments (gray diamonds) taken from Donovan et al., 2016a, and were collected in 2013 and 2014.

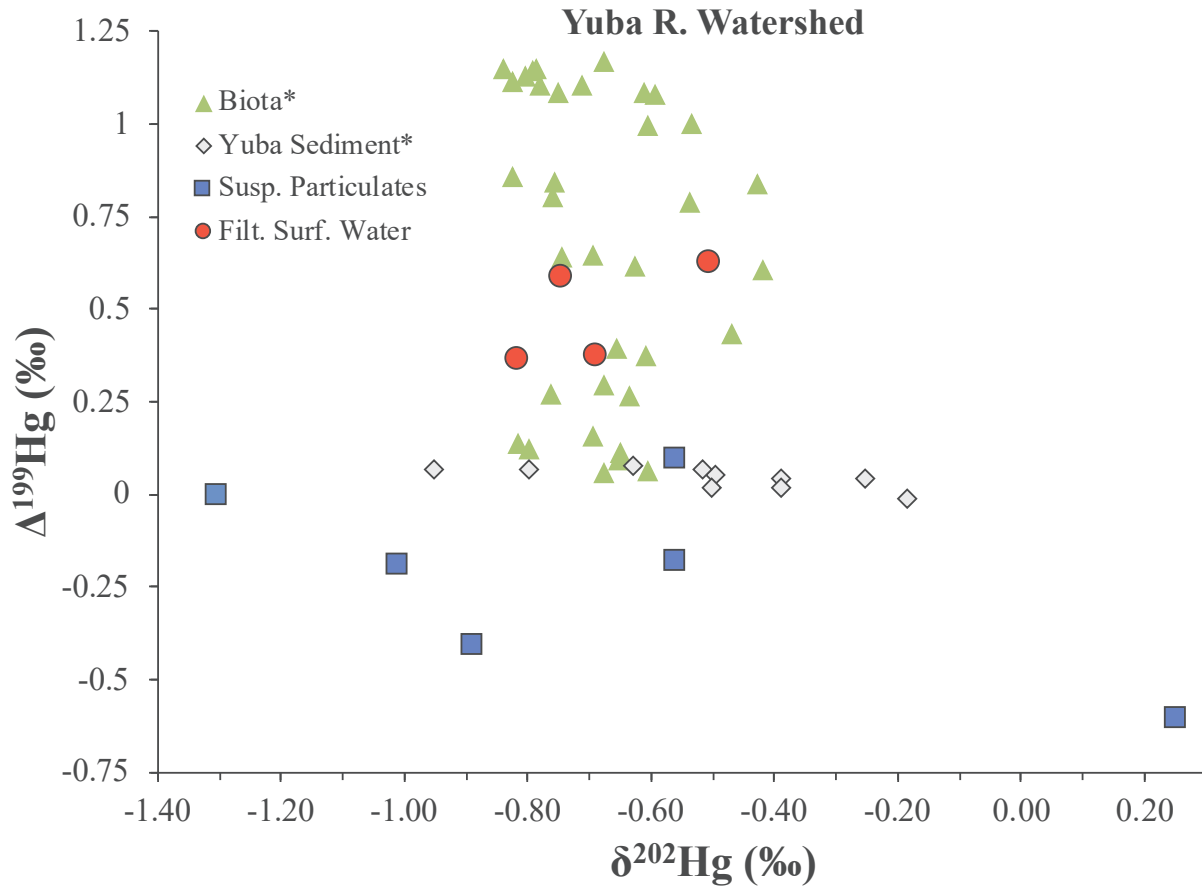


Figure 2

Plot of $\delta^{202}\text{Hg}$ (‰) vs $\Delta^{199}\text{Hg}$ (‰) values for samples from the Cache Creek watershed, including sites on Bear Creek and Sulfur Creek. Filtered surface waters (red circles) and suspended particulates (blue squares) were collected in June 2016, while values for biota (green triangles) and riverbed and terrace sediments (gray diamonds) taken from Donovan et al., 2016b, and were collected in 2013 and 2014. Note the difference in scales compared to Fig 1.

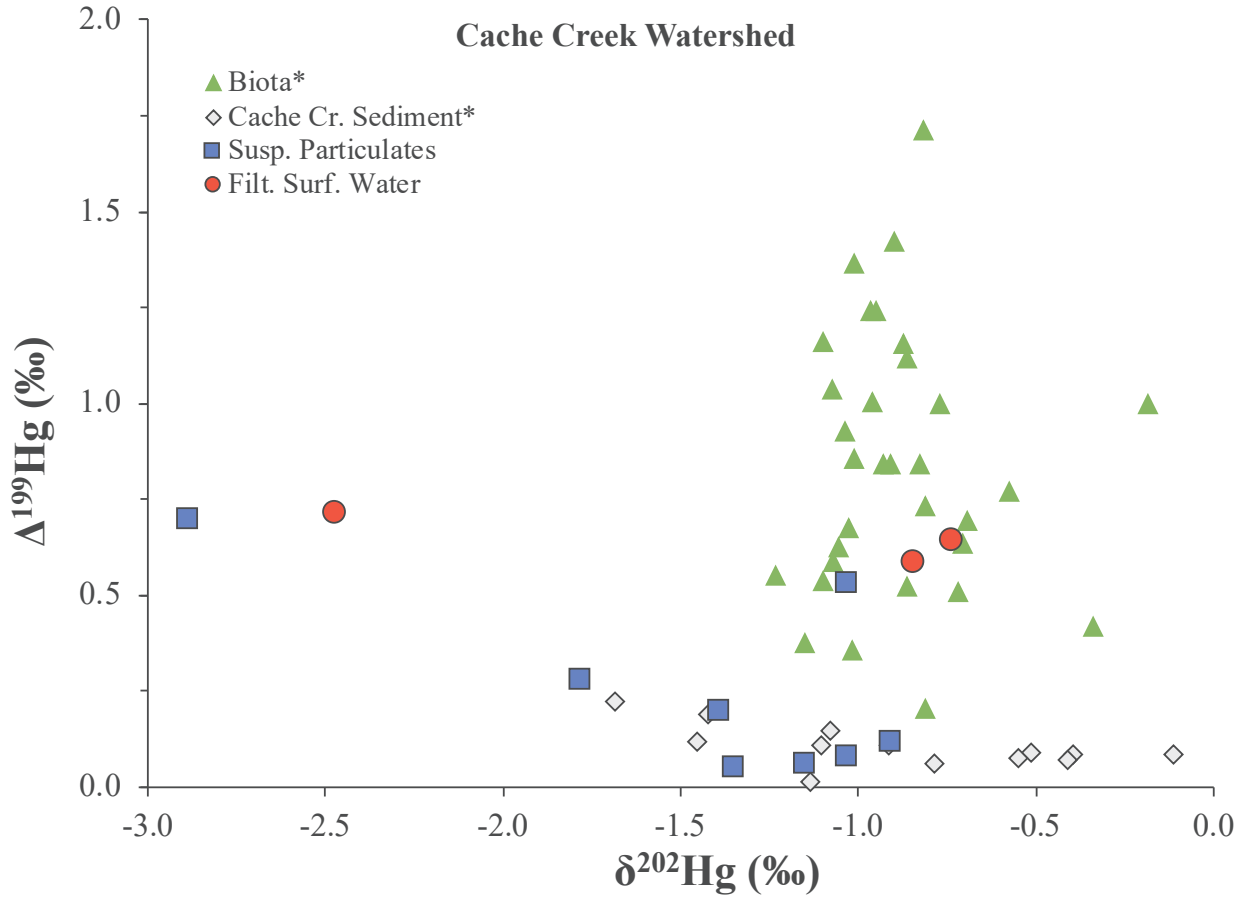


Figure 3

Plot of $\delta^{202}\text{Hg}$ (‰) vs $\Delta^{199}\text{Hg}$ (‰) values for paired riverbed and terrace sediments (diamonds), suspended particulates (squares), and filtered surface waters (circles). Samples from various sampling sites are denoted by the symbol color displayed in the figure legend. Sediment data taken from Donovan et al., 2016a and Donovan et al., 2016b.

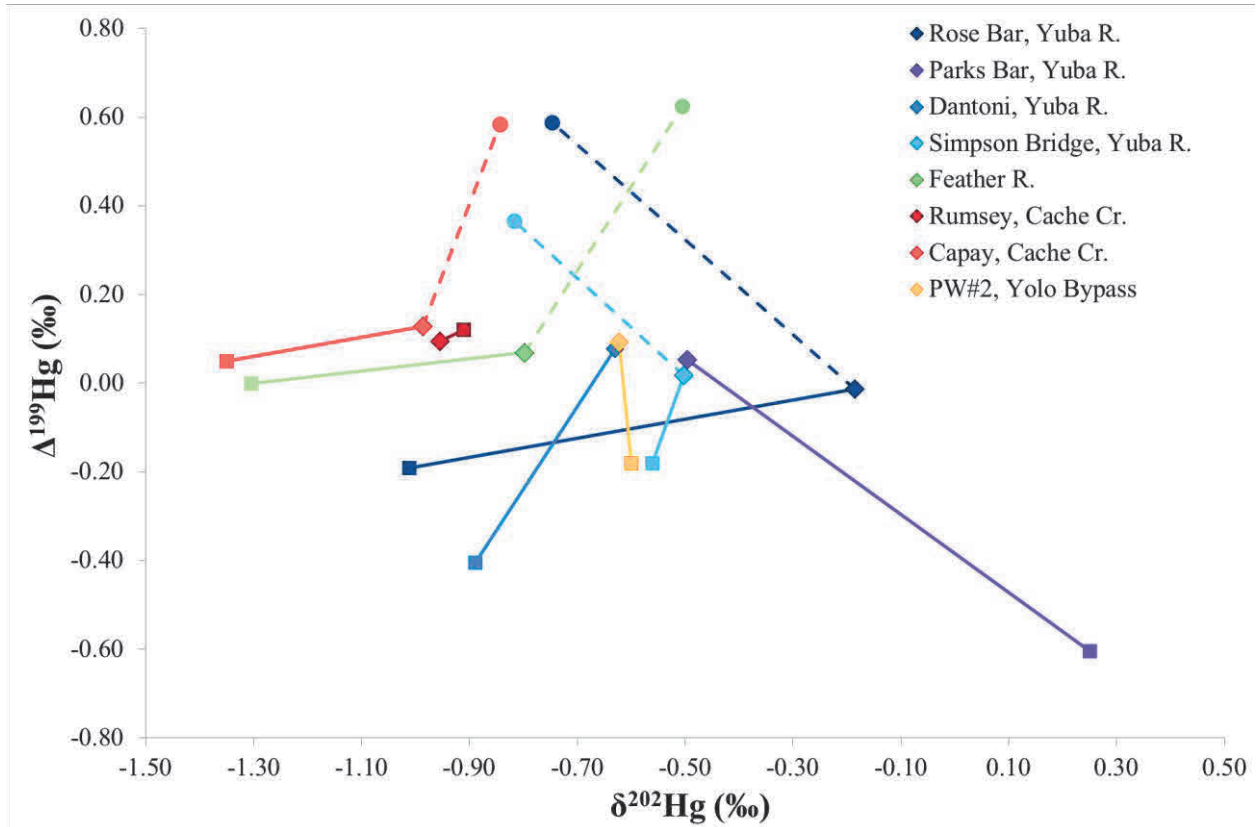


Figure 4

Plot of $\Delta^{201}\text{Hg}$ (‰) vs $\Delta^{199}\text{Hg}$ (‰) values for all filtered surface water samples (circles) and suspended particulates (squares) collected in both the Yuba River (red symbols), and Cache Creek (blue symbols) watersheds. The slope value for the combined watershed dataset (solid black line), only Cache Cr. dataset (solid blue line), and only Yuba R. dataset (solid red line), were calculated using a bivariate York regression (slope \pm 2SE [shown as dotted black lines for combined slope]), as described in the text in Section 3.2.1. The two samples denoted with blue asterisks had low recoveries that were unlikely to affect MIF signatures but were not included in the linear regression calculations, as described in Section 2.2.

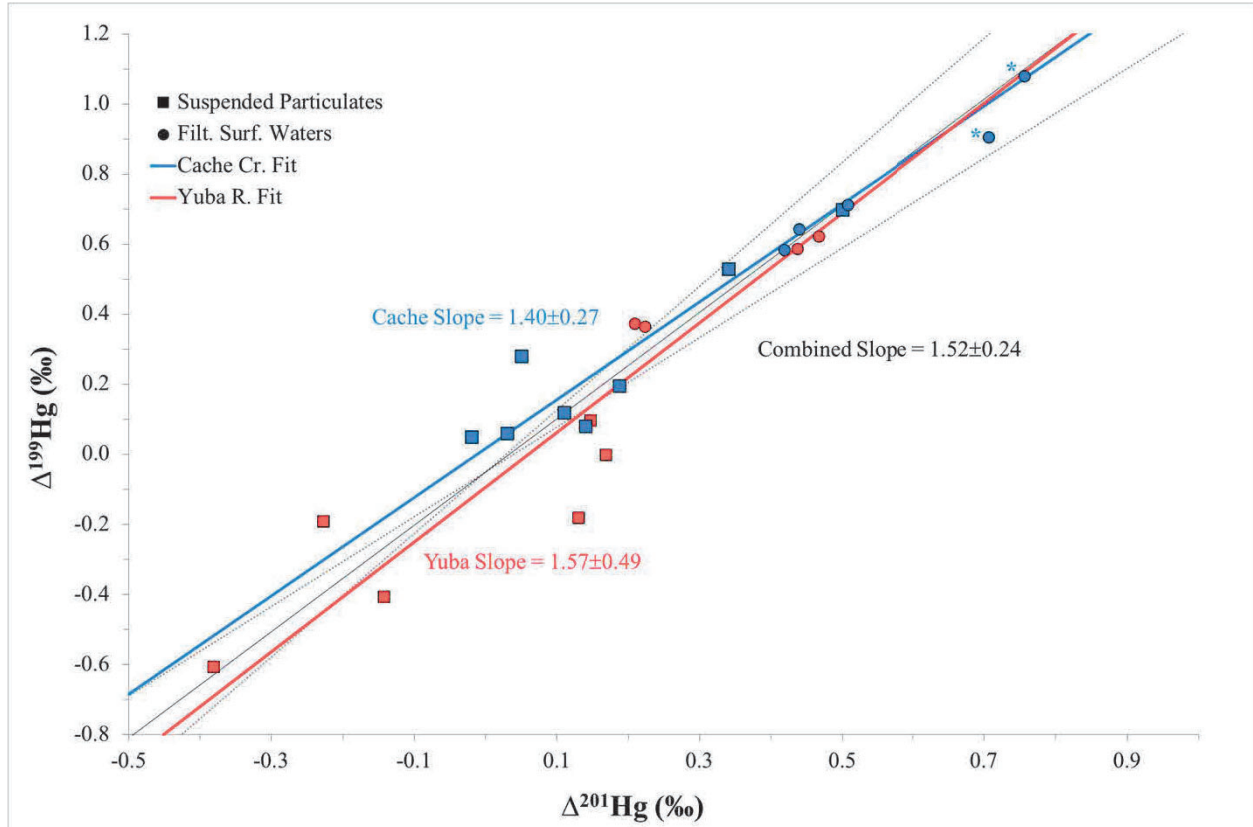


Table 1

Summary of THg concentration and Hg stable isotope data of collected samples from the CA Central Valley watersheds, including filtered surface waters (“FSW”) and the associated suspended particulates (“SW Susp. Part.”) from the Yuba R. and Cache Cr. watersheds and Yolo Bypass Wilderness Area. FSW samples denoted with ** (values highlighted in grey) had low recoveries that were unlikely to affect MIF signatures, but may have affected MDF signatures as discussed in Section 2.2

Watershed	Location	Sample Type	THg	THg	log(K _d)	δ ²⁰⁴ Hg	δ ²⁰² Hg	δ ²⁰¹ Hg	δ ²⁰⁰ Hg	δ ¹⁹⁹ Hg	Δ ²⁰⁴ Hg	Δ ²⁰¹ Hg	Δ ²⁰⁰ Hg	Δ ¹⁹⁹ Hg
			ng/L	ng/g		‰	‰	‰	‰	‰	‰	‰	‰	‰
Cache Creek	Bear Cr. Above Sulfur Cr.	SW Susp. Part.		1698.6	5.98	-2.48	-1.39	-0.86	-0.66	-0.15	-0.40	0.19	0.04	0.20
		FSW	1.77			-1.14	-0.73	-0.11	-0.33	0.46	-0.05	0.44	0.04	0.64
	Sulfur Cr.	SW Susp. Part.		57614.0	5.51	-4.39	-2.89	-1.67	-1.34	-0.03	-0.08	0.50	0.11	0.70
		FSW	179.08			-3.74	-2.47	-1.35	-1.18	0.09	-0.05	0.51	0.06	0.71
	Bear Cr. at Holston Chimney Canyon	SW Susp. Part.		2788.2	5.37	-1.52	-1.03	-0.43	-0.49	0.27	0.02	0.34	0.03	0.53
		**FSW	11.78			-2.28	-1.54	-0.40	-0.72	0.69	0.01	0.76	0.05	1.08
	North Fork Cache Cr.	SW Susp. Part.		210.7		-1.33	-1.03	-0.63	-0.52	-0.18	0.20	0.14	0.00	0.08
	Regional Park	SW Susp. Part.		114.6		-1.90	-1.15	-0.84	-0.57	-0.23	-0.18	0.03	0.01	0.06
	Rumsey	SW Susp. Part.		154.2	5.18	-1.44	-0.91	-0.57	-0.43	-0.11	-0.09	0.11	0.02	0.12
		**FSW	1.03			-1.59	-1.06	-0.09	-0.48	0.64	-0.02	0.71	0.05	0.91
Guinda	SW Susp. Part.		223.3		-2.59	-1.78	-1.29	-0.82	-0.17	0.07	0.05	0.08	0.28	
Capay	SW Susp. Part.		472.0	5.44	-1.81	-1.35	-1.03	-0.62	-0.29	0.20	-0.02	0.06	0.05	
	FSW	1.89			-1.28	-0.84	-0.22	-0.39	0.37	-0.02	0.42	0.03	0.58	
Yuba River	Rose Bar	SW Susp. Part.		166.1	5.65	-0.92	-1.01	-0.99	-0.55	-0.45	0.59	-0.23	-0.04	-0.19
		FSW	0.48			-1.12	-0.75	-0.12	-0.34	0.40	-0.01	0.44	0.03	0.59
	Parks Bar	SW Susp. Part.		665.2		0.66	0.25	-0.19	-0.17	-0.54	0.29	-0.38	-0.29	-0.61
	HammonGrove	SW Susp. Part.		401.1	5.72	-0.94	-0.56	-0.28	-0.26	-0.04	-0.10	0.15	0.02	0.10
		FSW	0.86			-1.09	-0.69	-0.31	-0.34	0.20	-0.06	0.21	0.01	0.37
	Dantoni	SW Susp. Part.		612.4		-0.50	-0.89	-0.81	-0.72	-0.63	0.83	-0.14	-0.27	-0.41
	Simpson Bridge	SW Susp. Part.		361.3	5.71	-1.13	-0.56	-0.29	-0.29	-0.32	-0.29	0.13	-0.01	-0.18
		FSW	0.74			-1.22	-0.82	-0.39	-0.40	0.16	0.00	0.22	0.01	0.37
	Feather R.	SW Susp. Part.		238.3	5.62	-1.78	-1.31	-0.81	-0.62	-0.33	0.17	0.17	0.03	0.00
		FSW	0.57			-0.79	-0.50	0.09	-0.22	0.50	-0.04	0.47	0.03	0.62
Yolo Bypass Wilderness Area	Permanent Wetlands #2	SW Susp. Part.		222.0	5.23	-0.86	-0.60	-0.53	-0.42	-0.33	0.03	-0.09	-0.12	-0.18
		FSW	1.30			NA								

Table 1

Summary of THg concentration and Hg stable isotope data of collected samples from the CA Central Valley watersheds, including filtered surface waters (“FSW”) and the associated suspended particulates (“SW Susp. Part.”) from the Yuba R. and Cache Cr. watersheds and Yolo Bypass Wilderness Area. FSW samples denoted with ** (values highlighted in grey) had low recoveries that were unlikely to affect MIF signatures, but may have affected MDF signatures as discussed in Section 2.2

Watershed	Location	Sample Type	THg ng/L	THg ng/g	log(K _d)	δ ²⁰⁴ Hg ‰	δ ²⁰² Hg ‰	δ ²⁰¹ Hg ‰	δ ²⁰⁰ Hg ‰	δ ¹⁹⁹ Hg ‰	Δ ²⁰⁴ Hg ‰	Δ ²⁰¹ Hg ‰	Δ ²⁰⁰ Hg ‰	Δ ¹⁹⁹ Hg ‰
Cache Creek	Bear Cr. Above Sulfur Cr.	SW Susp. Part.		1698.6	5.98	-2.48	-1.39	-0.86	-0.66	-0.15	-0.40	0.19	0.04	0.20
		FSW	1.77			-1.14	-0.73	-0.11	-0.33	0.46	-0.05	0.44	0.04	0.64
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		FSW	179.08			-3.74	-2.47	-1.35	-1.18	0.09	-0.05	0.51	0.06	0.71
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		FSW	0.48			-1.12	-0.75	-0.12	-0.34	0.40	-0.01	0.44	0.03	0.59
	Parks Bar	SW Susp. Part.		665.2		0.66	0.25	-0.19	-0.17	-0.54	0.29	-0.38	-0.29	-0.61
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		FSW	0.57			-0.79	-0.50	0.09	-0.22	0.50	-0.04	0.47	0.03	0.62
Yolo Bypass Wilderness Area	Permanent Wetlands #2	SW Susp. Part.		222.0	5.23	-0.86	-0.60	-0.53	-0.42	-0.33	0.03	-0.09	-0.12	-0.18
		FSW	1.30			NA								

Figure

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Yuba R. Watershed

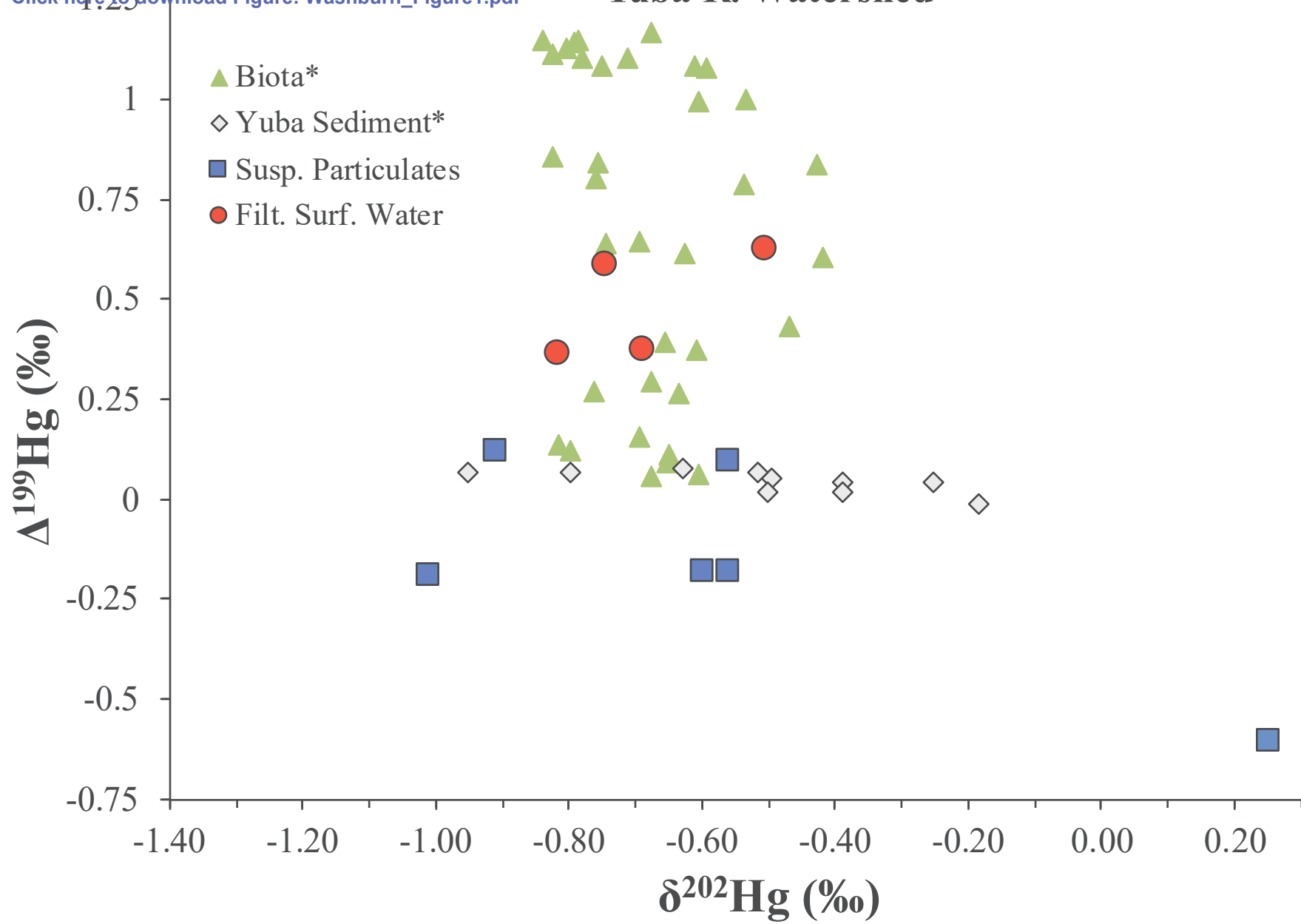


Figure 2.0

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Cache Creek Watershed

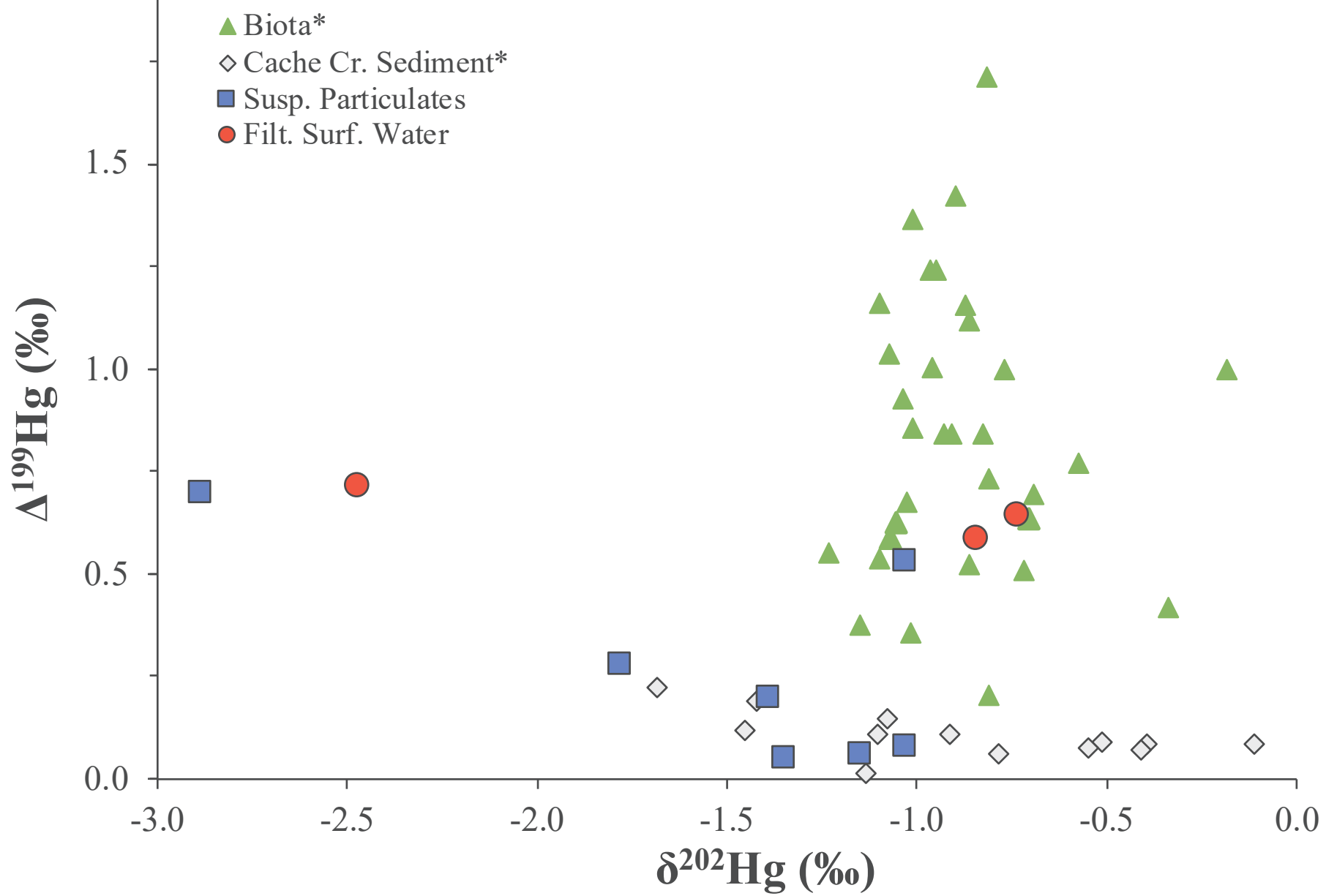


Figure
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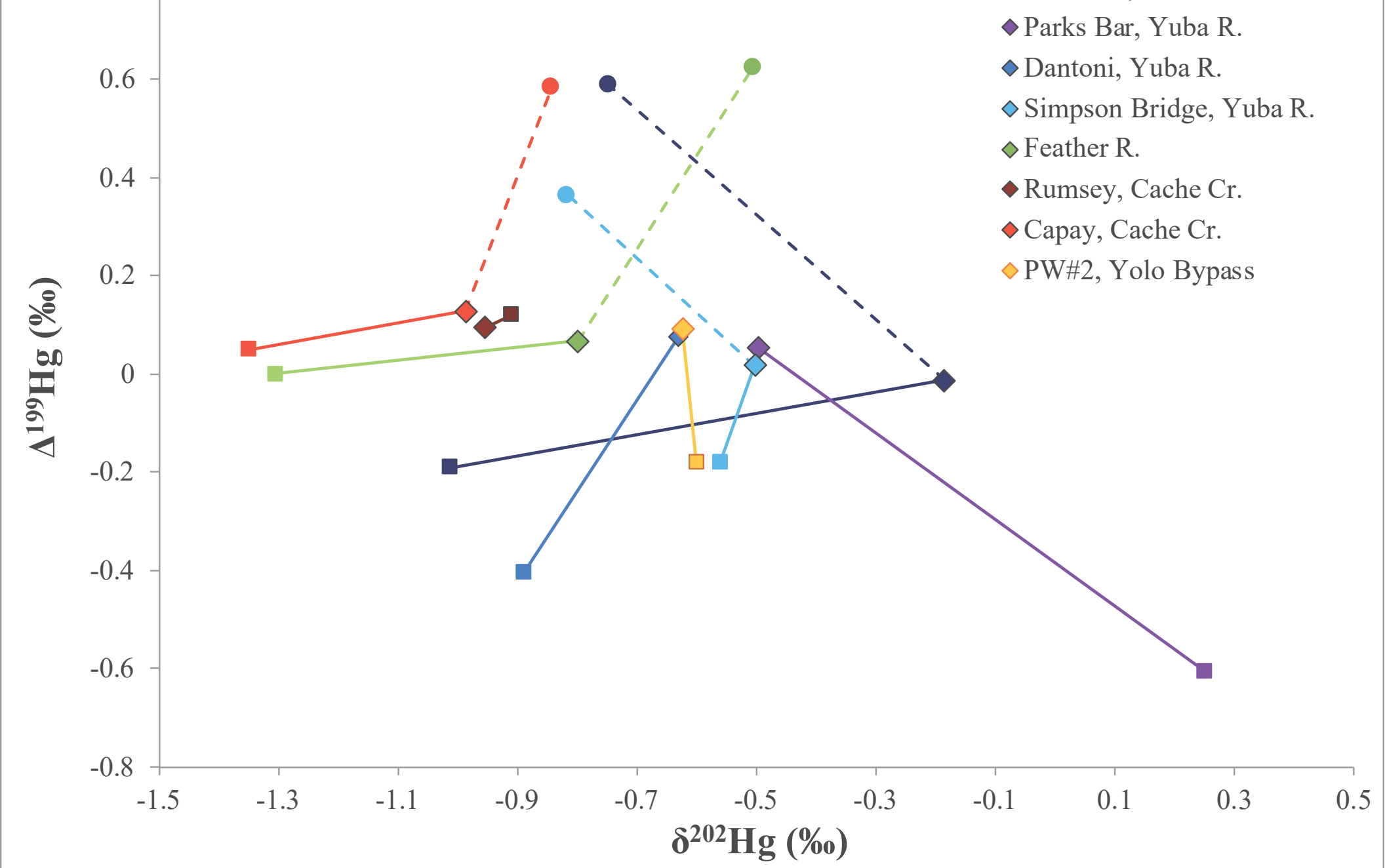
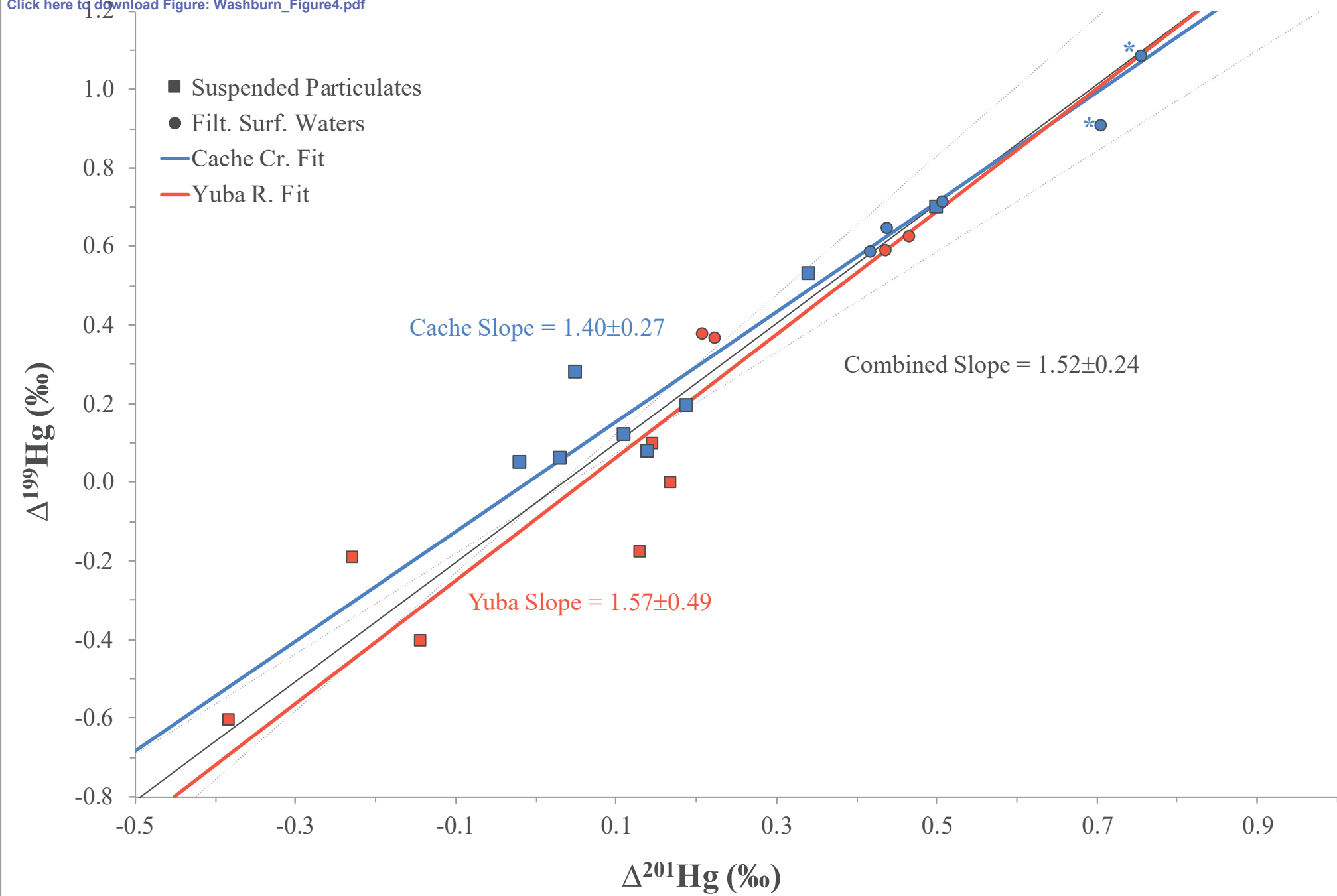


Figure
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