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# Understanding the mobilisation of metal pollution associated with historical mining in a carboniferous upland catchment

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## Abstract

Point and diffuse pollution from metal mining has led to severe environmental damage worldwide. Mine drainage is a significant problem for riverine ecosystems, it is commonly acidic (AMD), but neutral mine drainage (NMD) can also occur. A representative environment for studying metal pollution from NMD is provided by Carboniferous catchments characterised by a circumneutral pH and high concentrations of carbonates, supporting the formation of secondary metal-minerals as potential sinks of metals. The present study focuses on understanding the mobility of metal pollution associated with historical mining in a Carboniferous upland catchment. In the uplands of the UK, river water, sediments and spoil wastes were collected over a period of fourteen months, samples were chemically analysed to identify the main metal sources and their relationships with geological and hydrological factors. Correlation tests and principal component analysis suggest that the underlying limestone bedrock controls pH and weathering reactions. Significant metal concentrations from mining activities were measured for zinc (4.3 mg/l), and lead (0.3 mg/l), attributed to processes such as oxidation of mined ores (e.g. sphalerite, galena) or dissolution of precipitated secondary metal-minerals (e.g. cerussite, smithsonite). Zinc and lead mobility indicated strong dependence on biogeochemistry and hydrological conditions (e.g. pH and flow) at specific locations in the catchment. Annual loads of zinc and lead (2.9 and 0.2 tonnes/year) demonstrate a significant source of both metals to downstream river reaches. Metal pollution results in a large area of catchment having a depleted chemical status with likely effects on the aquatic ecology. This study provides an improved understanding of geological and hydrological processes controlling water chemistry, which is critical to assessing metal sources and mobilization, especially in neutral mine drainage areas.

## 39 **1. Introduction**

40 Anthropogenic activities have become an important driver of the global biogeochemical cycling  
41 of metals. Present day and historical mining have caused the release of heavy metals into  
42 fluvial environments. Globally, pollution from metal mining has led to severe damage to  
43 riverine ecosystems in many catchments<sup>1-6</sup>. In the United Kingdom, old mines from the 18th  
44 and 19th centuries represent the major diffuse source of metals having an adverse effect on  
45 aquatic ecosystems<sup>7-10</sup>. Rivers draining these mining areas are heavily affected by metal  
46 pollution as mineral veins present elevated concentrations of lead and zinc ores with variable  
47 concentrations of cadmium, barium and fluorine<sup>11</sup>.

48  
49 Water chemistry in surface waters are particularly vulnerable to biogeochemical and  
50 hydrological processes which are controlled by seasonality<sup>12-14</sup>. In this context, knowledge  
51 about metal mobility in natural water systems is extremely complex<sup>15</sup>. Biogeochemical  
52 partitioning of metals results in a diversity of forms. Within the dissolved phase metals are  
53 present as hydrated free ions, and associated with organic and inorganic complexes. Within  
54 the suspended particulate phase, metals may be complexed with inorganic or organic particles  
55 and biota or be present as discrete metal minerals. Adsorption and desorption of metals  
56 depend on a number of factors including pH, redox conditions, mineral ore sources and the  
57 composition of suspended particulate matter. For this reason, developing an improved  
58 understanding of the mechanisms determining the mobility and toxicity of metals within aquatic  
59 ecosystems is a key issue, which can in turn support efforts to manage or mitigate pollution  
60<sup>16</sup>.

61  
62 Several countries have developed guidelines to obtain good ecological and chemical status of  
63 and ground waters<sup>17</sup>. In the European Union (EU), the implementation of the Water  
64 Framework Directive (WFD) obliges member states to assess surface waters through  
65 improved catchment scale management (River Basin Management Plans, RBMPs). However,  
66 surface water bodies such as headwater streams have been excluded from early RBMPs due  
67 to their small size. Studies from Freeman et al.,<sup>18</sup> Dodds and Oakes,<sup>19</sup> and Meyer et al.<sup>20</sup> have  
68 shown the importance of these waterbodies as biodiversity richness, migration corridors, origin  
69 of stream networks and diffuse source of chemicals. Consequently, sound management is  
70 crucial for maintaining ecosystem health in higher order streams that are targeted by the WFD  
71 aims.

72  
73 A serious environmental hazard caused by mining is the generation of acid mine drainage  
74 (AMD). Mine drainages, spoil wastes run-off and spoil erosion constantly discharge large  
75 amounts of dissolved and particulate metals through AMD, representing a persistent and acute

76 pollution source and reducing water and sediment quality <sup>13, 14, 21</sup>. Studies of mine wastes  
77 chemistry have identified two types of mine effluents, acid mine drainage (low pH and high  
78 concentration of dissolved sulphate) and circumneutral mine drainage (major ion  
79 concentrations reflect the mineralogy of the catchment bedrock) <sup>22, 23</sup>. Thus, mine drainage is  
80 dependent on the geologic setting, local water chemistry, kinetic rates, and permeability of ore  
81 and gangue minerals <sup>24-28</sup>. Mine drainage flowing through Carboniferous limestone host rock  
82 is consequently metal-rich but with a circumneutral pH <sup>13, 14</sup>. Research from Lindsay et al.<sup>29</sup>  
83 and Desbarats and Dirom<sup>30</sup> indicates that circumneutral mine drainage might support natural  
84 attenuation of some metal-sulphides (e.g. ZnS, PbS) through the precipitation of secondary  
85 minerals. Consequently, catchments under these conditions may be more vulnerable to  
86 environmental harm due to changes in geochemical or hydrological conditions, producing high  
87 pulses of dissolved metal concentrations or long leaching processes in response to decades  
88 of chemical weathering <sup>31, 32</sup>.

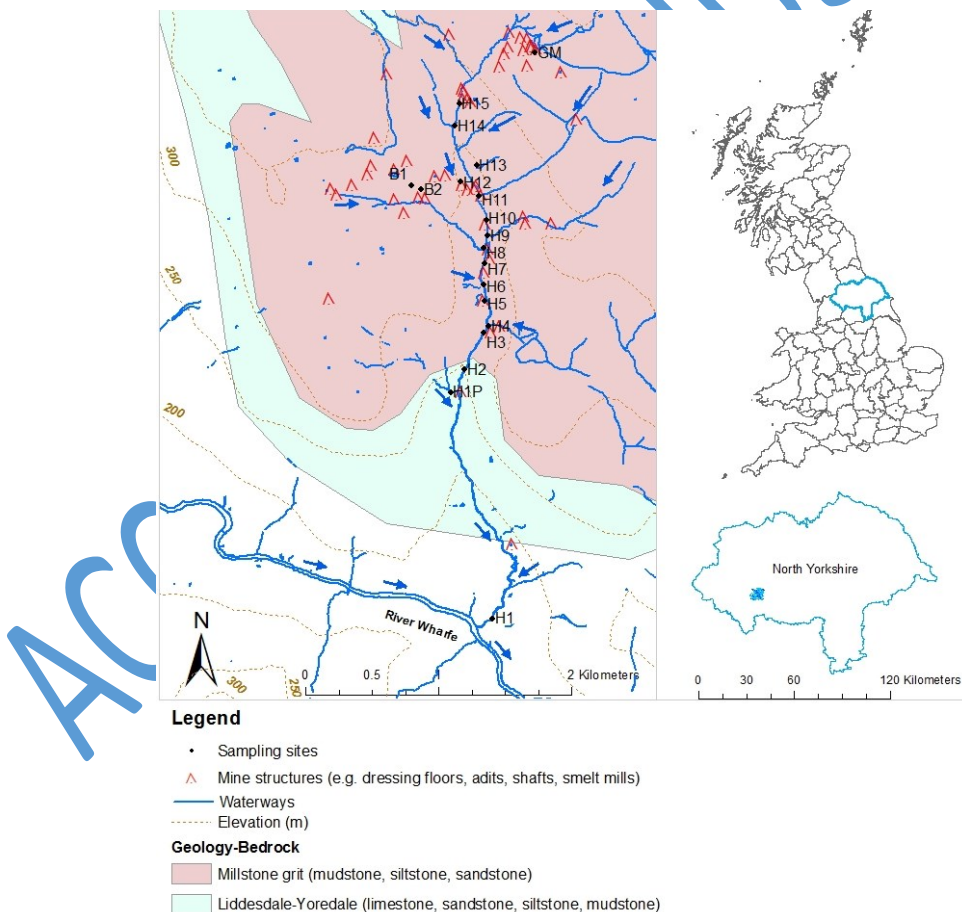
89  
90 Another long-standing metal pollution problem is physical and chemical mobilisation of metals  
91 through the passive dispersal and active transformation of abandoned tailings, spoil heaps,  
92 bed sediments and contaminated floodplains. Studies on metal transportation from mine  
93 wastes have reported the mobility of metals over long distances as free ions and complexed  
94 forms within rivers. In addition, solid phases can be stored within floodplain deposits for  
95 decades to millennia <sup>8, 33-37</sup>. In an area of the UK with Carboniferous bedrock, the north  
96 Pennines, historical metal mining has directly affected surface and subsurface floodplain soils  
97 with heavy metal concentrations above background levels <sup>38</sup>. Specifically, the Yorkshire Ouse  
98 basin which drains the Pennine Orefield is estimated to contain 620 million tonnes of lead and  
99 640 million tonnes of zinc stored within its floodplains <sup>3</sup>. Given the large differences in  
100 chemistry between acid and circumneutral mine drainage, particularly in the concentrations of  
101 protons and of Fe and Al whose solubility is controlled by pH, there will be significant  
102 differences in the degree of availability of metal forms that can interact with aquatic organisms.  
103 This necessitates dedicated studies of such Carboniferous catchments.

104  
105 This study aims to understand metal occurrence and mobilisation in a Carboniferous limestone  
106 upland catchment impacted by former lead and zinc mining. Comprehensive water monitoring  
107 and analysis of sediment and spoil samples are used to describe the effects of historical  
108 mining on a whole small river catchment in the northern Pennines region of North Yorkshire,  
109 UK. We sought to provide a better understanding of how geochemical processes control the  
110 concentration and mobility of dissolved metals in neutral metal-rich drainage. The results have  
111 wider implications for management strategies of potential environmental harm in such  
112 catchments.

113 **2. Site characteristics of Hebden Beck**

114 **2.1. Lithology and geology**

115 Hebden Beck is a sub-catchment of the River Wharfe located in the northern Pennines region  
116 of the United Kingdom and within the Yorkshire Dales National Park. Hebden Beck rises from  
117 Grassington Moor and is joined by multiple tributaries including Coalgrove Beck, Bolton Gill  
118 and Loss Gill before the confluence with the Wharfe. It is approximately 12 km long, with the  
119 upper reach (6.4 km) of the main channel being the most heavily impacted by historic mine  
120 working. It drains an area of 26 km<sup>2</sup> and subsequently flows into the River Wharfe which drains  
121 the Wharfedale valley. The Wharfe flows into the River Ouse and ultimately reaches the sea  
122 at the Humber Estuary which is one of the largest in the UK (24,750 km<sup>2</sup>). The geology of  
123 Hebden Beck is dominated by Millstone Grit sandstone (approx. 20 km<sup>2</sup>) but crucially also  
124 contains bands of Carboniferous limestone (approx. 6 km<sup>2</sup>) (Figure 1). The catchment cover  
125 comprises 46% peatlands, predominantly in the upstream areas, 35% Carboniferous  
126 limestone, mainly in the south and 19% glacial sediment, predominantly in the west<sup>39</sup>.



127

128 Figure 1. Hebden Beck with sampling sites and mine structures located in the Millstone grit and  
129 Liddesdale-Yoredale bedrock. Blue arrows indicate the direction of flow.

130 2.2. Ore processing and mine wastes

131 Hebden's orefield comprised coal and lead-zinc mineral deposits, where galena (PbS) is the  
132 most common mineral, but with associated sphalerite (ZnS), chalcopyrite (CuFeS<sub>2</sub>), barite  
133 (BaSO<sub>4</sub>), fluorite (CaF<sub>2</sub>) calcite (CaCO<sub>3</sub>) and witherite (BaCO<sub>3</sub>)<sup>39</sup>. These minerals occur mostly  
134 in vertical veins along fault planes. Early mine workings were open cuts and shafts to extract  
135 deeper layers of lead ore. In later years until around 1850, the hushing method was used to  
136 scour away the soil using the erosive power of water to expose mineral veins. This method  
137 required the construction of dams to control streams, and manmade channels to divert water.  
138 Horizontal drainage levels (adits) were driven from the valley bottoms to enable deeper  
139 working and easier removal of minerals. Lead ores were crushed, classified and bagged at  
140 the dressing floors located at the surface close to the mines, then transported to the smelting  
141 mills to be processed. Water power was also applied at the dressing floors and smelt mills,  
142 therefore spoil tips or mine wastes are located next to rivers. Approximately, 124 mining  
143 features exist in the Hebden Beck catchment area, however main features are 5 lead-zinc  
144 mines, 15 adits, 7 spoil tips and 4 smelters. From 1700-1900, 1686.5 tonnes of lead were  
145 extracted from these mines<sup>40</sup>.

146 **3. Methods**

147 3.1. Sampling strategy

148 Sites were selected from the most impacted area, covering an area of 5 km<sup>2</sup> including point  
149 and diffuse sources flowing downstream from mine sites. A total of sixteen sites were chosen  
150 for water sampling including the main channel, minor and major tributaries, together with a  
151 source pool feeding a tributary (Table 1). Eight of these sites (about 3.3 km) were part of a  
152 water quality monitoring programme performed by the Environment Agency<sup>39</sup>. The other  
153 eleven sites were selected based on their proximity to mine wastes (e.g. tailings, spoils).  
154 Monthly sampling campaigns were carried out from November 2013 to December 2014.

155 Table 1. Sampling sites along Hebden Beck. Table indicates type of sample, site elevation, coordinates,  
156 and distance from River Wharfe. Sites are listed from upstream to downstream. Locations adopted from  
157 the Environment Agency monitoring programme are indicated with (\*).

Site ID	Site Description	Type of sample	Elev. (m)	Coordinates		From R. Wharfe (m)
				East	North	
H15	Head water reservoir (Next-smelt mill)	Water	368	402798	466766	4970
GM	Grassington moor, spoil wastes	Spoil	380	403014	466663	
H14	Perennial tributary (Coalgrove Beck)	water, sediment	294	402413	466106	4207
H13	Ephemeral tributary	Water	287	402443	465931	4030
B1	Beaver, spoil wastes (from heap)	Spoil	320	402087	465660	
B2	Beaver, spoil wastes (silt runoff)	Spoil	317	402163	465630	
H12	Ephemeral tributary (downstream-Yarnbury mine-Beaver spoil)	water, sediment	285	402451	465822	3921

<b>H11</b>	Perennial tributary (Loss Gill Dike)	Water	278	402597	465578	3637
<b>H10*</b>	Main channel	water, sediment	267	402656	465324	3377
<b>H9*</b>	Perennial tributary (Bolton Gill)	Water	266	402661	465285	3338
<b>H8*</b>	Ephemeral tributary (from-Yarnbury mine)	Water	268	402632	465176	3226
<b>H7*</b>	Perennial tributary (Adit) – Bolton Haw	Water	266	402648	465164	3206
<b>H6*</b>	Main channel	Water	257	402630	464916	2958
<b>H5*</b>	Perennial tributary (Duke's adit)	Water	256	402638	464793	2836
<b>H4</b>	Ephemeral tributary (Waterfall)	Water	254	402668	464604	2645
<b>H3*</b>	Perennial tributary (Laneshaw adit)	Water	246	402632	464550	2580
<b>H2*</b>	Main channel at gauging station	Water	235	402488	464275	2271
<b>H1P</b>	Perennial tributary	water	233	402382	464104	2071
<b>H1</b>	Main channel - Confluence R. Wharfe	water, sediment	152	402695	462400	140

158

### 159 3.2. Water sampling

160 Samples were taken from downstream to upstream (H1 to H15) in order to minimise  
161 contamination of other sites by disturbance<sup>41</sup>. At each site, a sample was taken with a pre acid  
162 washed (10% HNO<sub>3</sub>, Nitric acid-Sigma Aldrich 69% and Milli-Q water) 750 ml polypropylene  
163 bottle attached to a plastic pole. Four subsamples were then extracted from this bottle. For  
164 total metals, unfiltered samples were placed individually into a pre-weighted 50 ml tube  
165 (polypropylene) containing 1 ml of preservation solution (10% HNO<sub>3</sub>) to reach 1% v/v of the  
166 final volume and pH ≤ 2<sup>41, 42</sup>. For dissolved metals analysis, samples filtered through syringe  
167 filters (0.45 µm, polyethersulfone-hydrophilic, Sartorius) were placed individually into a pre-  
168 weighted 50 ml tube (polypropylene). Then preservation solution (10% HNO<sub>3</sub>) was added as  
169 used for total metals. For quantifying major anions, the sample was filtered (Sartorius syringe  
170 filters 0.45 µm, polyethersulfone-hydrophilic) and placed into polypropylene tubes. For  
171 inorganic and organic carbon analysis the samples were passed through syringe filters (0.45  
172 µm, nylon-polypropylene, Avonchem) and placed into polypropylene tubes. All samples were  
173 kept in a cool box during sampling and transported the same day to the laboratory for storage.  
174 Samples for major anions analysis were stored frozen at -20°C, while samples for all other  
175 analyses were refrigerated at 4°C.

176 A carbon Analyser (Analytik Jena Multi N/C2100) was used for measuring carbon compounds  
177 (dissolved inorganic carbon-DIC and dissolved organic carbon-DOC), Ion Chromatographer  
178 for major cations (Ca, Mg) and anions (Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup>) (Dionex ICS-3000), and SEAL  
179 Analytical AA3 was used for orthophosphate quantification. For metal analysis, nine elements  
180 were measured (Pb, Ba, Cd, Sr, Zn, Cu, Fe, Mn, Al), using inductively coupled plasma mass  
181 spectrometry (ICP-MS; Thermo Fisher iCAPQc) with specific limits of detection (Pb: 0.0001  
182 µg/l, Ba: 0.06 µg/l, Cd: 0.0001 µg/l, Sr: 0.08 µg/l, Zn: 67 µg/l, Cu: 0.05 µg/l, Fe: 0.11 µg/l, Mn:  
183 0.04 µg/l, Al: 0.16 µg/l). Field blanks (n= 3) and replicates (n= 3) were collected at each  
184 sampling campaign.

### 185 3.3. Sediment and spoil sampling

186 Most sites were dominated by large rocks and coarse sediment. Sediment samples were  
187 collected during a single campaign at specific sites (H14, H12, H10 and H1) to assess the  
188 evolution of mineral composition. Plastic scoops were used for their collection by wading along  
189 a cross-section of the stream. Sediments were sieved through a <250 µm stainless steel mesh  
190 and transferred into 50ml polypropylene tubes. Spoil sampling was carried out at single spoil  
191 heaps at Grassington Moor (GM) and the Beaver spoil area (B1), in addition a further sample  
192 was collected at the Beaver spoil area that represented material that had been subjected to  
193 movement and size sorting by the actions of rainfall events (B2).

194 In the laboratory, sediment samples were centrifuged at 3200 rpm for 10 mins to allow removal  
195 of the supernatant. The supernatant was decanted and the resulting slurries were placed in a  
196 petri dish to air dry. After drying, sediments were placed into zip log bags for disaggregation.  
197 Spoil samples (B1, B2, and GM) were dried to calculate percentage water composition. The  
198 surface area was also measured to estimate the mineral area available for dissolution  
199 reactions. This was performed using the Brunauer, Emmett and Teller method (BET;  
200 Micromeritics Gemini VII 2390a) on 2 g of sample dried overnight under N<sub>2</sub> gas at 75 °C. In  
201 addition, both sediment and spoil samples were analysed by X-ray Diffraction (XRD, Bruker  
202 D8-Discover instrument) for determining mineralogy of the major constituents and X-ray  
203 Fluorescence (XRF, Innovex X-5000) for chemical composition. The minimum mineral  
204 fractions required for detection on this instrument is 2-3%. The standard reference material  
205 STSD-3 (stream sediment) was used as XRF quality control.

### 206 3.4. In situ measurements

207 Pre-calibrated multiple sensor probes (Model HQ30d flexi 1032) were used in the field to  
208 measure pH, dissolved oxygen (DO: mg/l) and conductivity (EC: µS/cm). Flow rate (m<sup>3</sup>/s) was  
209 calculated from in situ flow velocity measurements (m/s) (flow meter: Global 800-876) together  
210 with data from river depth (m) and width (m). Flow data from the UK Environment Agency  
211 gauging station (H2) was also obtained from their continuous monitoring records. This flow  
212 data together with metal concentrations were used for the calculation of annual metal loading  
213 and comparison with Environmental quality standards (EQS) for freshwater in the UK<sup>39</sup>.

### 214 3.5. Data analysis

215 Geochemical modelling

216 The PHREEQC code (version 3)<sup>43, 44</sup> was used for modelling main geochemical reactions  
217 occurring in aqueous solutions. This software allows the prediction of mineral precipitation that  
218 potentially controls the composition of the aqueous phase. Equilibrium reactions and  
219 thermodynamic constants were retrieved from the built-in WATEQ4F database<sup>45-47</sup>. Mineral

220 saturation indices and metal free ion activities for hydroxide, carbonate and sulphate minerals  
 221 were calculated for the pH range 3.5-9 and based on mean values across our field sites and  
 222 all sampling dates: temperature 10 °C, SO<sub>4</sub><sup>2-</sup> ( $\bar{x}$ : 13592 µg/l) and Cl<sup>-</sup> ( $\bar{x}$ : 7730 µg/l). With  
 223 calculations for carbonate minerals the pCO<sub>2</sub> was fixed at three times the atmospheric  
 224 concentration (0.0012 atm), consistent with typical supersaturation of this gas in streams.  
 225 These model predictions are compared with metal free ion activities calculated for each  
 226 sampling site and date to investigate the controlling mineral phases.

## 227 Principal Component Analysis

228 Principal component analysis (PCA) was conducted to identify the main factors influencing  
 229 metal distribution. Linear correlation analysis was applied to evaluate the relationships among  
 230 the studied metals, other compounds and in situ parameters. Results of Pearson and  
 231 Spearman tests showed no significant difference between them. As such, we report the  
 232 Pearson correlation, as this test is more sensitive for the identification of outliers. Both test  
 233 were performed using Rstudio (version 3.1.0).

## 234 4. Results

### 235 4.1. Characterising metal pollution in the catchment

236 In water samples, metal concentrations occurred in the following order for total:  
 237 Zn>Fe>Sr>Ba>Pb>Al>Mn>Cd>Cu and dissolved forms: Zn>Fe>Sr>Ba>Al>Pb>Mn>Cd>Cu  
 238 (Table 1-SI). Two metals, Zn and Pb were chosen as the focus for this study based on their  
 239 significant concentrations derived from mining activities (Table 2). Annual pH averages  
 240 reflected the considerable contribution of the underlying limestone bedrock showing a  
 241 dominant circumneutral pH (mean= 6.8) in 80% of the studied sites. Some sites (e.g. H4 and  
 242 H13) represent moorland runoff with little interaction with underlying rock, thus pH is lower  
 243 (<6.3) due to high DOC and no carbonate buffering.

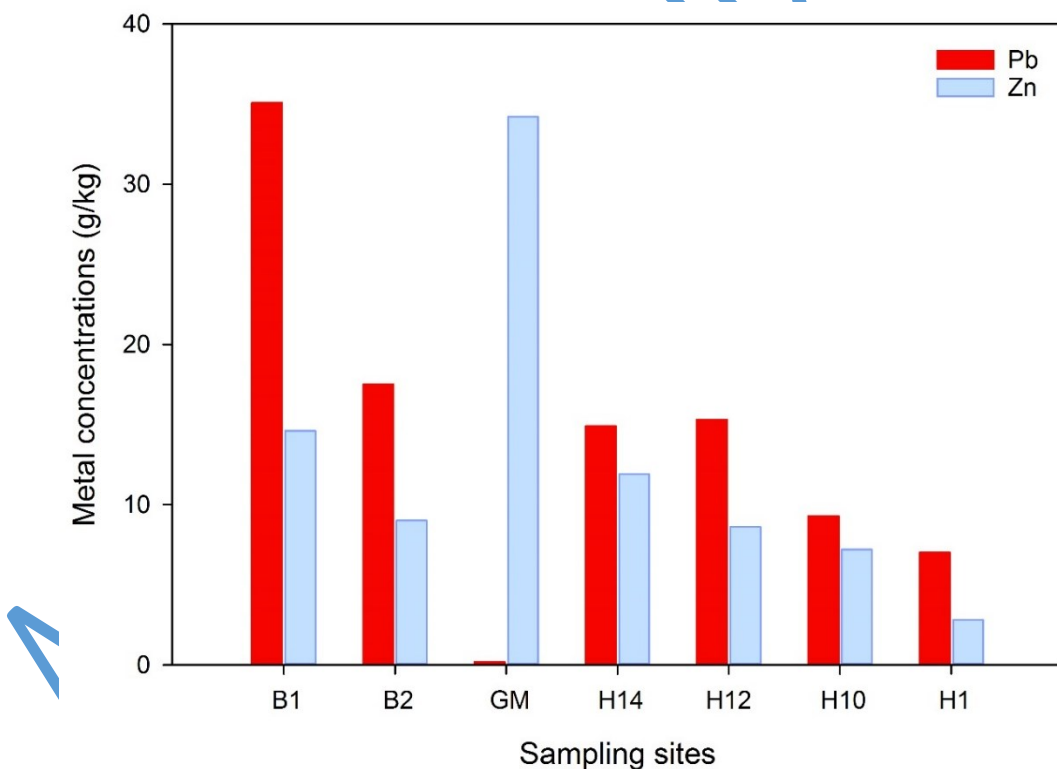
244 Table 2. Mean, maxima and minima from water chemical analysis. Metal forms are denoted as total (T)  
 245 and dissolved (D). Description of sites are indicated as main channel (MC), ephemeral tributaries (ET)  
 246 and perennial tributaries (PT). Units are in µg/l. Values below the detection limit are represented by  
 247 (b/d).

Site	Description		Pb <sub>T</sub>	Pb <sub>D</sub>	Zn <sub>T</sub>	Zn <sub>D</sub>
H15	Reservoir	Ave	316.7	279.6	2058.8	2028.0
		Max	411.9	423.6	2542.4	2759.5
		Min	96.3	103.6	722.4	1080.7
H14	PT	Ave	178.3	157.6	1318.6	1397.9
		Max	292.3	375.5	1778.9	2193.1
		Min	106.9	75.7	864.7	900.3
H13	ET	Ave	227.1	205.9	260.4	207.4
		Max	331.3	283.4	1619.8	1435.5
		Min	141.2	138.0	b/d	b/d
H12	ET	Ave	686.4	284.2	5168.8	4252.3

		Max	2701.2	439.9	12619.3	7438.4
		Min	261.8	184.1	66.5	73.6
H11	PT	Ave	64.0	31.6	295.7	410.5
		Max	765.6	355.2	3493.9	5276.2
		Min	3.7	b/d	b/d	b/d
H10	MC	Ave	108.1	80.7	468.8	444.4
		Max	268.1	145.2	787.4	777.0
		Min	11.8	3.1	b/d	b/d 33.5
H9	PT	Ave	28.5	20.8	125.2	95.7
		Max	171.9	94.3	572.2	518.6
		Min	12.7	2.0	b/d	b/d
H8	ET	Ave	38.1	19.0	318.5	269.0
		Max	123.5	28.2	438.3	435.2
		Min	12.2	5.3	b/d	b/d
H7	PT	Ave	7.7	4.3	3440.2	3220.5
		Max	21.1	47.7	5425.8	4312.3
		Min	0.8	b/d	2062.0	1936.7
H6	MC	Ave	60.4	49.4	537.6	510.2
		Max	102.0	85.6	664.1	674.1
		Min	20.5	13.8	468.3	390.2
H5	PT	Ave	60.4	54.3	883.5	867.2
		Max	132.3	157.8	1216.5	1206.0
		Min	17.1	10.0	688.0	613.5
H4	ET	Ave	8.3	4.4	b/d	b/d
		Max	26.7	6.6	68.6	68.6
		Min	3.2	b/d	b/d	b/d
H3	PT	Ave	2.8	0.2	b/d	b/d
		Max	10.6	0.7	98.6	84.7
		Min	0.05	b/d	b/d	b/d
H2	MC	Ave	46.7	39.4	515.0	485.9
		Max	93.5	87.4	765.4	606.9
		Min	17.0	4.2	194.5	360.5
H1P	PT	Ave	3.0	1.9	b/d	b/d
		Max	5.2	8.3	68.6	68.5
		Min	0.8	b/d	b/d	b/d
H1	MC	Ave	31.1	16.6	217.5	158.9
		Max	146.8	43.7	375.9	302.9
		Min	5.1	b/d	77.9	67.8

248 Across the catchment, Zn<sub>D</sub> was the most abundant pollutant with concentrations ranging from  
249 95.7 to 3220.5 µg/l in perennial tributaries. Ephemeral tributaries also showed high  
250 concentrations up to 4252.3 µg/l while sites along the main channel had Zn<sub>D</sub> concentrations  
251 from 158.9 to 510.2 µg/l. The second toxic pollutant of concern was Pb<sub>D</sub>, where main  
252 contributions were observed in ephemeral tributaries with ranges from 4.4 to 284.2 µg/l, and  
253 perennial tributaries ranged from 1.9 to 157.6 µg/l. The main river channel showed  
254 concentrations of Pb from 16.6 to 80.7 µg/l. Major cations were dominated by calcium, with  
255 concentrations from 2.6-54.9 mg/l, and major anions comprised sulphate (3-24.9 mg/l), nitrate  
256 (1-17.8 mg/l), phosphate (0.002 to 0.1 mg/l) and chloride (7.2-10.5 mg/l). Dissolved inorganic  
257 carbon concentrations ranged from 0.9 to 42.7 mg/l and dissolved organic carbon from 1.2 to  
258 16.8 mg/l (Table 2-SI). Field blanks measurements shown concentrations below limit for all  
259 the elements and replicates a standard deviation of ≤ ±0.5 µg/l (Cd and Cu), ≤ ±3.6 µg/l (Mn  
260 and Sr), ≤ ±12 µg/l (Pb, Al, Ba).

261 From mineralogical analysis of spoil and sediment samples, the most abundant minerals were  
 262 quartz ( $\text{SiO}_2$ ) and fluorite ( $\text{CaF}_2$ ). Spoil samples (B1, B2 and GM) included barite ( $\text{BaSO}_4$ ) as  
 263 an additional dominant mineral. Other secondary minerals like muscovite  
 264 ( $\text{KAl}_2(\text{Si}_3\text{AlO}_{10})(\text{OH})_2$ ) and kaolinite ( $\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$ ) were present at B1 and GM. Furthermore,  
 265 the mineral cerussite ( $\text{PbCO}_3$ ) was detected at B1. In sediments (H14, H12, H10, H1), the  
 266 mineralogy of H14 was similar to B2 (principally the presence of quartz, fluorite and barite)  
 267 whereas H12 and H10 contained calcite ( $\text{CaCO}_3$ ), and H1 (140 m from the confluence with  
 268 the River Wharfe) presented mainly quartz and calcite. Chemical composition analysis  
 269 detected significant fractions of Pb in spoils B1 (35.1 g/kg) and Zn in GM (34.2 g/kg), while for  
 270 sediments, major Pb concentrations were present in H12 (15.3 g/kg) and Zn in H14 (11.9  
 271 g/kg). Total Pb plus Zn in the sediment samples showed a decrease the further downstream  
 272 the sample origin (Figure 2). Analysis of water composition and surface area in spoil samples  
 273 showed higher percentages of water composition in GM (34.29%) followed by B2 (18.77%)  
 274 and B1 (18.6%), while surface area values were: GM (20.6  $\text{m}^2/\text{g}$ ), B1 (5.82  $\text{m}^2/\text{g}$ ), B2 (1.34  
 275  $\text{m}^2/\text{g}$ ).



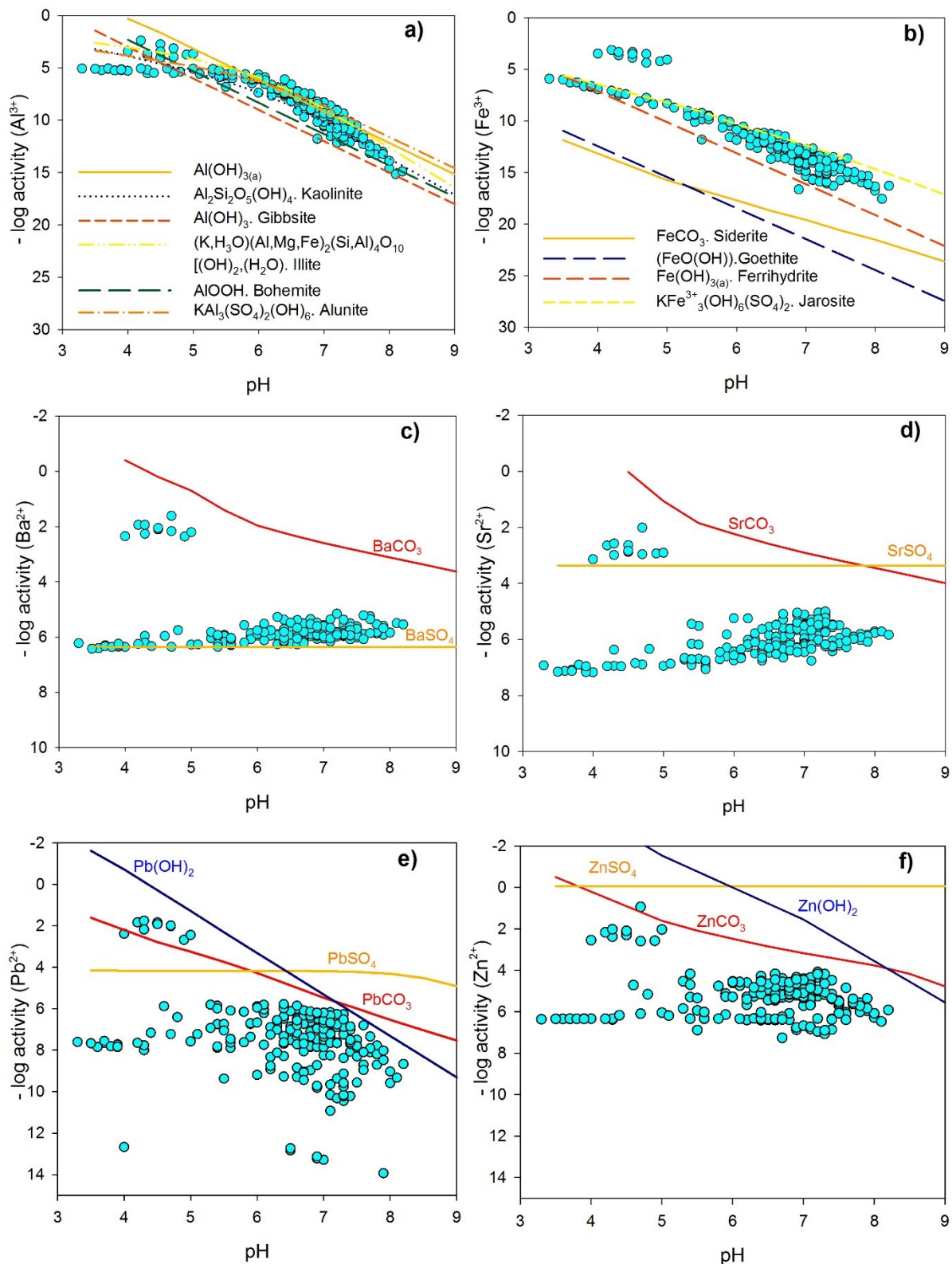
276  
 277 Figure 2. Lead and zinc composition in spoils and sediments.

278 **4.2. Mineral phases controlling dissolved metal concentrations**

279 The metals Zn and Pb are the most significant toxic pollutants derived from mining activities.  
 280 However, we modelled the geochemical behaviour of the additional metals Al, Fe, Ba and Sr  
 281 as they are present in significant concentrations in the catchment. Geochemical modelling

282 predicted that kaolinite ( $\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$ ) may account for the Al source with the solubility being  
283 controlled mainly by amorphous Al oxide phases ( $\text{Al}(\text{OH})_3$ ). Concentrations of Fe are more  
284 likely derived from secondary minerals. Iron solubility is controlled by amorphous phases such  
285 as ferrihydrite ( $\text{Fe}(\text{OH})_3$ ). The supersaturation of the Al and Fe phases may suggest the  
286 presence of some colloidal metal measured as part of the dissolved fraction. Barium  
287 concentrations are controlled by barite ( $\text{BaSO}_4$ ), while Sr activity was too low to infer a  
288 controlling phase, perhaps being controlled by a mineral where it is present as a secondary  
289 metal. Lead and Zn concentrations are largely regulated by secondary minerals as metal-  
290 carbonates, cerussite ( $\text{PbCO}_3$ ) and smithsonite ( $\text{ZnCO}_3$ ), respectively (Figure 3).

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292 Figure 3. Aluminium, Fe, Ba, Sr, Pb and Zn activity as a function of pH,  $\text{SO}_4^{2-}$  (13592  $\mu\text{g/l}$ ),  $\text{Cl}^-$  (7730  
 293  $\mu\text{g/l}$ ) and  $p\text{CO}_2 = 0.0012$  atm. Theoretical saturation of mineral forms are represented by solid lines  
 294 and calculated metal free ion activity of experimental data by dots.

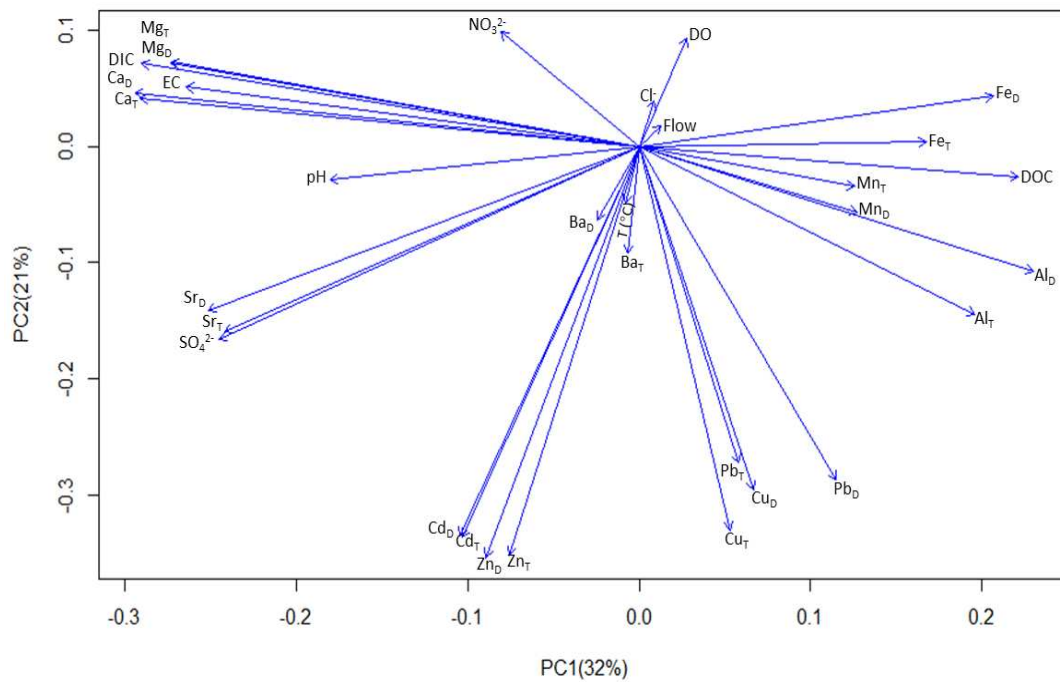
295 4.3. Key chemical relationships

296 Table 3 summarises correlation analysis of metals derived from mining activities (Pb, Zn) and  
 297 bedrock weathering (Ca) with major water chemistry parameters (pH, SO<sub>4</sub><sup>2-</sup>, DIC and DOC).  
 298 Zinc presented good correlation with SO<sub>4</sub><sup>2-</sup> (r= 0.6), Ca showed very strong association with  
 299 DIC (r= 0.9), while Pb presented poor correlations (r≤ 0.3). Principal component analysis  
 300 (PCA) shows key geochemical processes in influencing the water chemistry of Hebden Beck  
 301 (Figure 4). The first component (PC1) with a 32% of variance indicates strong correlations  
 302 between dissolved and particulate forms of calcium, magnesium, with DIC, EC and pH,  
 303 reflecting weathering of the bedrock, while moderate correlation between iron and DOC refers  
 304 to the transport of metals through colloidal matter in aquatic systems. Furthermore, the second  
 305 component (PC2) with 21% of variance shows good correlations between strontium, zinc and  
 306 cadmium with SO<sub>4</sub><sup>2-</sup>, reflecting the oxidation of sulphide minerals. Other in situ parameters like  
 307 temperature, DO, flow and anions (NO<sub>3</sub><sup>-</sup> and Cl<sup>-</sup>) were not significantly associated with metals  
 308 when considering all sites.

309 Table 3. Relationships between metals (Pb, Zn, and Ca) with pH, SO<sub>4</sub><sup>2-</sup>, DIC and DOC. Pearson  
 310 correlation coefficient is denoted as r, p-value as p and confidence interval (95%) as CI.

	Stats	pH	SO <sub>4</sub> <sup>2-</sup>	DIC	DOC
<b>Pb<sub>D</sub></b>	r	-0.1	-0.1	-0.4	0.3
	p	0.084	0.194	<0.001	<0.001
	CI	[-0.257 0.016]	[-0.228 0.047]	[-0.517 -0.284]	[0.135 0.393]
<b>Zn<sub>D</sub></b>	r	0.2	0.6	0.04	-0.2
	p	0.008	<0.001	0.588	0.022
	CI	[0.051 0.319]	[0.517 0.692]	[-0.101 0.176]	[-0.295 -0.024]
<b>Ca<sub>D</sub></b>	r	0.6	0.7	0.9	-0.6
	p	<0.001	<0.001	<0.001	<0.001
	CI	[0.526 0.698]	[0.571 0.730]	[0.963 0.978]	[-0.701 -0.529]

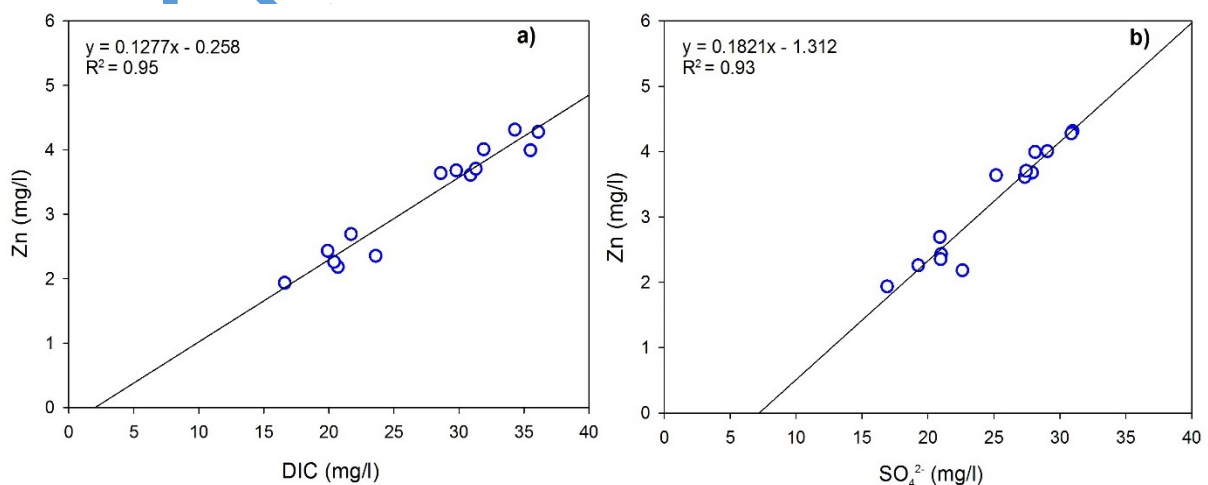
311



312

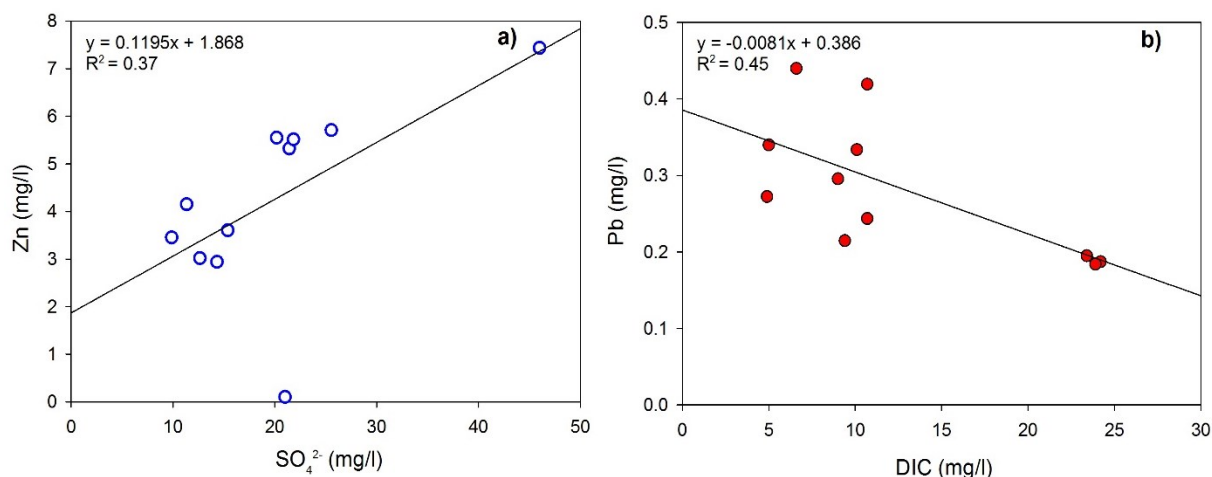
313 Figure 4. Scree plot from PCA analysis, horizontal axis shows projections of the first principal  
 314 component PC1 which represents 32% of the total variance and the vertical axis the second component  
 315 PC2 representing the 21% of variance.

316 Complementary linear regression analysis was carried out at sites with highest concentrations  
 317 of  $Zn_D$  and  $Pb_D$  (H15, H14, H12 and H7) for evaluating their relationships with DIC and  $SO_4^{2-}$ .  
 318 Since H15 is a reservoir with lower variance in chemical concentrations, it was not considered  
 319 for this and subsequent analyses. Strong correlations between  $Zn_D$  and DIC ( $R^2= 0.95$ ) and  
 320  $SO_4^{2-}$  ( $R^2= 0.93$ ) were identified at H7. Moderate correlations between  $Zn_D$  and  $SO_4^{2-}$  ( $R^2= 0.4$ )  
 321 and between Pb and DIC ( $R^2= 0.5$ ) were present at site H12 (Figure 5 and 6). No significant  
 322 correlation was evident at site H14, therefore results are not shown.



323

324 Figure 5. Relationships in tributary H7 between  $Zn_D$  and DIC (left panel) and  $SO_4^{2-}$  (right panel).



325

326 Figure 6. Relationships in tributary H12 between  $\text{Zn}_D$  with  $\text{SO}_4^{2-}$  and  $\text{Pb}_D$  with DIC.

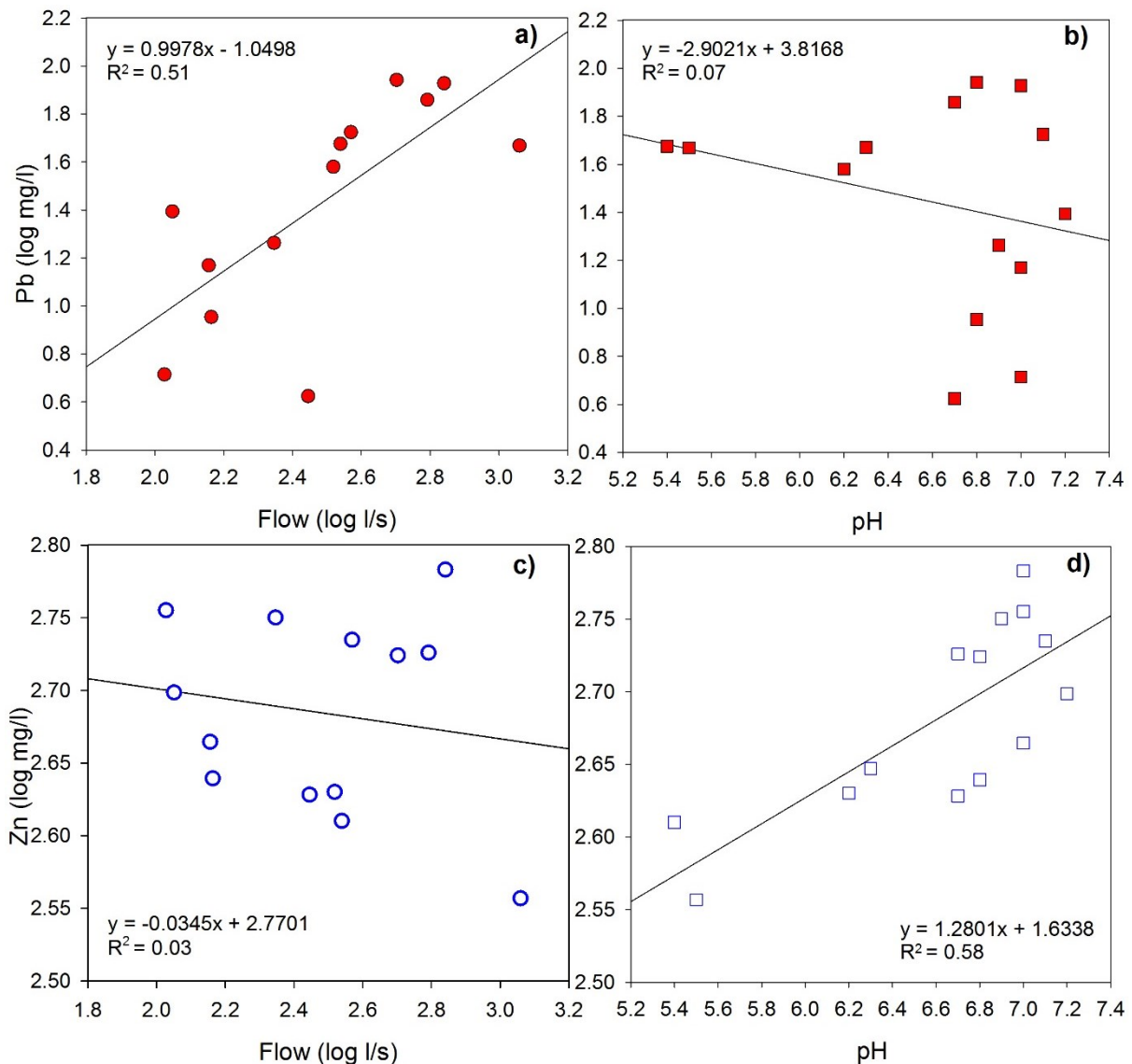
#### 327 4.4. Seasonality and trends

328 Across the sampling campaigns, water temperature ranged from 3.6 to 15.6 °C, with higher  
 329 values in summer (July) and lowest in winter (February). Consistent with temperature, low  
 330 monthly average of EC values were recorded in winter (78  $\mu\text{S}/\text{cm}$ ) and high average values  
 331 during summer (209  $\mu\text{S}/\text{cm}$ ). Dissolved oxygen levels showed similar values, ranging from  
 332 10.2-14.0 mg/l, with lower concentrations recorded in winter (December) and highest levels in  
 333 autumn (September). For flow, highest annual rates were recorded in November (2666.1 l/s)  
 334 and lowest rates in July (54.0 l/s) (Table 3-SI).

335 Fluctuations were also dependent on the type of tributaries (e.g. perennial and ephemeral)  
 336 (Table 4-SI). Flow values in perennial tributaries ranged from 3.0 to 192.7 l/s and in ephemeral  
 337 from 9.0 to 57.0 l/s when flow was measurable, as some tributaries were dry for 5 or 6 months.  
 338 Across all tributaries, two significant sites were identified as metal contributors of  $\text{Zn}_D$  (H12:  
 339 4252.3  $\mu\text{g}/\text{l}$  and H7: 3220.5  $\mu\text{g}/\text{l}$ ) and  $\text{Pb}_D$  (H12: 284.2  $\mu\text{g}/\text{l}$ ) (Table 1). Across all sites, mean  
 340 values of pH were highest in September (7.1) and lowest in November 2013 (5.6) (Table 3-  
 341 SI). In individual sites, pH means ranged from 3.9 to 7.5, showing large monthly variations  
 342 ( $\text{SD} \geq \pm 0.7$ ) in sites H15, H11, H9, and H1P. Despite these variances a circumneutral pH (6.2-  
 343 7.4) was predominant in the catchment.

344 Trends of flow and pH were considered to analyse their influence on metals concentration.  
 345 Regression analysis was performed using all sampling sites. However, the results  
 346 demonstrate the site specific nature of trends, with no catchment wide trends revealed (Figure  
 347 2-SI). We investigate further the trends for H2, as the chemistry is representative of a  
 348 significant distance of the downstream reach and it has limited dilution before entering the

349 River Wharfe. Strong positive relationships were present for Pb<sub>D</sub>-flow (R<sup>2</sup>= 0.51) and Zn<sub>D</sub>-pH  
 350 (R<sup>2</sup>= 0.58) and no clear relationships for Pb<sub>D</sub>-pH (R<sup>2</sup>= 0.06) and Zn<sub>D</sub>-flow (R<sup>2</sup>= 0.02) (Figure  
 351 7).



352  
 353 Figure 7. Trends of metals in function of flow and pH in H2. Panel a and b show trends of Pb and panel  
 354 c and d indicate Zn trends. Solid lines represent regression lines.

#### 355 4.5. Annual metal load

356 The contribution of Pb<sub>D</sub> and Zn<sub>D</sub> in the catchment was estimated through the annual metal  
 357 load (tonne/year). Only site H2 was considered for this calculation due to the availability of  
 358 flow data and the lack of significant additional mine runoff downstream. Table 4 shows the  
 359 average annual load of Pb is 0.2 tonne/year and for Zn 2.9 tonne/year. These values were  
 360 compared with well-established EQS metals and showed maximum exceedances of 12-fold  
 361 for Pb and Zn.

362

363 Table 4. Estimation of annual Pb and Zn load by using flow records from Environment Agency (EA)  
 364 gauging station at H2 (main channel).

Stats	H2-Flow (l/s) from EA station	Pb <sub>D</sub> (µg/l)	Pb <sub>D</sub> (tonne/year)	Zn <sub>D</sub> (µg/l)	Zn <sub>D</sub> (tonne/year)
<b>Average</b>	189	39.4	0.2	485.9	2.9
<b>Maximum<sup>(a)</sup></b>	556	87.4	1.5	606.9	10.0
<b>Minimum<sup>(b)</sup></b>	36	4.2	<0.1	360.5	<0.9
<b><i>EQS-Hardness based</i></b>		<b>7.2</b>		<b>50.0</b>	

365 a) Maximum values recorded in February 2014.

366 b) Minimum values recorded in July 2014.

367

## 368 5. Discussion

369 In the catchment, biogeochemical actions such as weathering and erosion are significant  
 370 processes in the generation of dissolved metals, which are likely derived from bedrock  
 371 weathering and oxidation or dissolution of mineral ores. Carboniferous limestone bedrock,  
 372 mainly composed of calcite (CaCO<sub>3</sub>) is weathered, releasing significant amounts of calcium  
 373 and carbonate, and creating a neutralizing capacity and circumneutral pH in environments  
 374 surrounded by sulphide ore wastes. The influence of the geology in the catchment was shown  
 375 by strong relationships between Ca<sub>D</sub> with DIC ( $r = 0.9, p < 0.001$ ) and pH ( $r = 0.6, p < 0.001$ ). For  
 376 metals derived from mining activities, a potential primary source of Zn<sub>D</sub> is the oxidation of  
 377 sphalerite (ZnS) as a good correlation was observed between Zn<sub>D</sub> and SO<sub>4</sub><sup>2-</sup> ( $r = 0.6, p < 0.001$ )  
 378 (Table 3). However, additional Zn<sub>D</sub> concentrations may be attributed to the presence of  
 379 secondary zinc minerals (e.g. smithsonite)<sup>48</sup>. Contributions of Pb<sub>D</sub> are associated with  
 380 dissolution of metal-carbonate compounds (cerussite) rather than oxidation of metal-sulphide  
 381 as no correlation between Pb<sub>D</sub> with SO<sub>4</sub><sup>2-</sup> was identified ( $r \leq -0.1, p < 0.001$ ) (Table 3). The  
 382 presence of secondary minerals such as metal-carbonate might affect the solubility and  
 383 mobility of metals as they present slower dissolution kinetics than primary minerals<sup>49-52</sup>.

384

385 Geochemical modelling has revealed the importance of secondary minerals such as  
 386 carbonates, sulphates and hydroxides in the control of dissolved metals (Figure 3). For  
 387 instance, Pb and Zn concentrations are greatly influenced by the dissolution of metal-  
 388 carbonate forms (e.g. cerussite and smithsonite)<sup>49, 53</sup>. Carbonates released from the  
 389 dissolution of metal-carbonate also contribute to the river alkalinity, enhancing the buffering  
 390 capacity of the system. In addition, the source of Al can be associated with the presence of  
 391 kaolinite, which was identified by the XRD analysis in most of the spoils/sediments. The  
 392 presence of Fe concentrations can be associated with jarosite, and although this mineral was  
 393 not detected by XRD analysis, it is a common secondary Fe mineral in mining areas<sup>27, 28</sup>.  
 394 Figure 3 shows that Fe activity follows closely the prediction for jarosite, which at pH > 3 tends

395 to dissolve and release sulphate ions and  $\text{Fe}^{3+}$ <sup>54</sup>. In alkaline environments the activity of  $\text{Fe}^{3+}$   
396 is likely controlled by hydrous ferrous oxides, the presence of colloids in the dissolved fraction  
397 may explain the supersaturation. Aluminium and Fe solubility are controlled by amorphous  
398 phases, specifically  $\text{Al}(\text{OH})_3$  and  $\text{Fe}(\text{OH})_3$  (known as ferrihydrite). The presence of hydroxide  
399 compounds in the catchment could affect metal mobility, as they might sorb or co-precipitate  
400 with metals like Pb and Zn, acting as natural scavengers of these toxic elements<sup>55</sup>. Nordstrom  
401<sup>56</sup> indicated that hydrology is another factor influencing metals mobility as concentrations of  
402 constituents in natural waters depend to a large extent on the rate of dissolution relative to  
403 flow rate. This condition was observed in the upper site of the catchment (a pond) as  
404 concentrations of metals were closer to saturation due to longer residence time of the water  
405 than under stream flow conditions (Figure 1-SI). Clustered sites with high metal free ion  
406 activities were identified and associated with sites with low pH values (<5), possible caused  
407 by high DOC concentrations (>8 mg/l), which are not considered in the modelling. The  
408 presence of metal-organic complexes will reduce the activities of metal free ions.

409  
410 Analysis of sediments and spoils were consistent with water chemistry results showing Zn  
411 (11.9 and 34.2 g/kg) and Pb (15.3 and 35.1 g/kg) as major metals present in spoils due to  
412 lower extraction efficiency methods common in historical mining<sup>40</sup>. Preliminary sequential  
413 batch leaching experiments of spoil samples into deionised water showed consistent Zn and  
414 Pb concentrations of at least one order of magnitude above other metals across several  
415 leaching cycles (except Pb in GM where it is not present in significant concentrations). For  
416 leaching with acid (0.1 M HCl) most Zn was solubilised in the first two batches. Similar  
417 concentrations of Pb and Ca were leached but over five acid addition cycles, suggesting  
418 different dissociation kinetics of Pb and Zn minerals<sup>57</sup>. Future work will further explore the  
419 kinetics of leaching from the mineral forms in the spoils and sediments. Mineralogical results  
420 revealed the presence of secondary minerals such as cerussite ( $\text{PbCO}_3$ ) as a source of Pb.  
421 Current chemical characterisation is in agreement with previous studies of water quality, metal  
422 composition and flux in the Yorkshire Pennine Orefield<sup>14</sup>.

423  
424 The impact of former metal mining on water quality has been evidenced by Pb, Zn and Cd  
425 pollution in the Yorkshire Pennine region, particularly in Hebden Beck<sup>14, 39</sup>. In expanding the  
426 range of Hebden Beck tributaries from earlier works, we have included sites close to mine  
427 wastes (e.g. tailings, spoils) for the identification of principal sources of metals. Major  
428 contributions of  $\text{Zn}_D$  and  $\text{Pb}_D$  were identified from mine water discharges (H7) and spoil wastes  
429 (H12)<sup>14</sup>. For these two sites there are statistically significant relationships between Zn and Pb  
430 with  $\text{SO}_4^{2-}$  and DIC inferring the composition of their respective mineral sources. At site H12,  
431 an extended area covered by spoil wastes from Yarnbury mines, moderate correlations were

432 identified between  $Zn_D$  with  $SO_4^{2-}$  ( $R^2= 0.37$ ), and  $Pb_D$  with DIC ( $R^2= 0.45$ ) reflecting the  $Zn_D$   
433 contribution from the oxidation of sphalerite and  $Pb_D$  contribution from the dissolution of  
434 cerussite (Figure 6). These correlations revealed the type and grade of ores mined during the  
435 eighteen and nineteenth centuries, producing spoils with different particle sizes and  
436 permeabilities, influencing their capacity to form secondary minerals<sup>58</sup>. At site H7, a mine  
437 channel from Bolton Haw, strong correlations were observed between Zn and  $SO_4^{2-}$  ( $R^2= 0.93$ )  
438 and Zn and DIC ( $R^2= 0.95$ ) (Figure 5). Both correlations suggest the oxidation of sphalerite as  
439 the main Zn source, however, if all sulphate was from ZnS the expected molar ratio between  
440  $Zn_D$  and  $SO_4^{2-}$  should be 1:1 instead of the observed value of 5:1. Furthermore, the  
441 mineralogical and geochemical results did not show extensive evidence of pyrite mixed in  
442 mineral veins (correlation of  $Fe/SO_4^{2-}$   $r= -0.5$ ;  $p= 0.076$ ). Likewise, Dunham and Wilson<sup>59</sup> have  
443 reported that although pyrite, marcasite ( $FeS_2$ ) and bravoite ( $(Fe, Ni, Co)S_2$ ) are abundant in  
444 the orefield, they are found in minor amounts in veins, representing small quantities in relation  
445 with adjacent deposits rich in Fe discharges. Thus, secondary zinc minerals such as  
446 smithsonite ( $ZnCO_3$ ), hydrozincite ( $5ZnO \cdot 2CO_2 \cdot 3H_2O$ ) and hercynite  
447 ( $Zn_4[Si_2O_7](OH)_2 \cdot H_2O$ ), should be considered as possible sinks<sup>49</sup>. No correlation for Pb was  
448 observed in this site, probable due to the presence of low concentrations ( $4.3 \mu g/l$ ) as  
449 consequence of aging<sup>60</sup> or the sorption effect of biofilms, becoming a significant sink for Pb  
450<sup>61</sup>.

451  
452 In Hebden Beck, flow events can alter the river water chemistry and metal concentrations<sup>14</sup>.  
453 During base flow conditions, the circumneutral pH and buffering capacity are maintained by  
454 groundwater rather than surface water. This condition contributes to the presence of  
455 secondary zinc minerals and other carbonate minerals that sequester zinc, also influences the  
456 complexation of Pb with carbonate and organic matter, and affects its transformation to other  
457 forms like hydroxide, oxyhydroxide, hydroxysulfate minerals, limiting solubility and further  
458 weathering<sup>49, 52, 56, 62</sup>. Flow fluctuations caused by drought or heavy rainfall allowed the  
459 identification of major point and diffuse sources (H7 and H12) and their metal contributions  
460 under different flow events. At the point source H7 (mine adit-Bolton Haw), metal  
461 concentrations were generally constant at both flow conditions, therefore this site can be  
462 considered as a continuous source of metals (particularly for Zn:  $3220.5 \mu g/l$ ). At the diffuse  
463 source H12 (ephemeral tributary-draining spoil wastes from Yarnbury mine), metal  
464 concentrations become more significant during high flow, but greater contributions were also  
465 observed after dry periods (e.g. July [0 l/s,  $Zn_D$  and  $Pb_D$  below detection limits], August [8.4  
466 l/s,  $5709.0 \mu g Zn_D/l$ ,  $419.2 \mu g Pb_D/l$ ]). This might be explained by the capacity of soluble  
467 sulphate minerals to store metals (e.g. Zn) during dry seasons and release them into the

468 environment during wet seasons<sup>63</sup>. In addition, Byrne et al.<sup>64</sup> and Cánovas et al.<sup>65</sup> have  
469 indicated the influence of runoff produced by storms in increasing metals dissolved from  
470 weathered metal salts (smithsonite, cerussite) located in superficial mine spoils. Rothwell et  
471 al.<sup>66</sup> showed differences in metal concentrations not only between base and high flow  
472 conditions but also within and between storm events. Thus, further studies of metal  
473 concentrations and fluxes under a range of hydrological conditions are pertinent since the  
474 frequency and magnitude of floods are increasing the transport of dissolved and particulate  
475 metal forms from sources to river channels and floodplain soils, which are often used for  
476 agriculture<sup>67</sup>.

477 Seasonal variations of pH and flow were considered to assess metal mobility. In the main  
478 channel (H2) strong correspondence occurred between pH-Zn<sub>D</sub> (r= 0.7) and flow- Pb<sub>D</sub> (r= 0.6)  
479 while relationships for Pb<sub>D</sub>-pH and Zn<sub>D</sub>-flow were unclear. The absence of a relationship of Zn  
480 with flow suggests that dissolution of zinc minerals is not kinetically limited (Figure 7), although  
481 solubility has been shown to depend on mineral composition in some cases<sup>58</sup>. The Pb  
482 relationship with flow may be related to greater flushing of areas where minerals have had  
483 longer to leach Pb into waters (e.g. H15 or the ephemeral pond feeding H12) (Figure 2-SI).  
484 Sims et al.<sup>68</sup> have also reported the role of flow in the generation of suspended matter,  
485 affecting the transport Pb forms. Once they enter into the aquatic system they tend to be  
486 adsorbed to suspended matter, while for the case of carbonate minerals they are likely to  
487 break down in acid waters, liberating significant quantities of Pb to sediments further down the  
488 river<sup>69</sup>. Thus, understanding the chemical tendencies of Zn and Pb under local pH and flow  
489 conditions is extremely important for estimating the potential fate and extent of pollutants.

490 Metal contributions from point and diffuse sources decreased downstream (H2) (Zn<sub>D</sub>: 479.4  
491 µg/l, Pb<sub>D</sub>: 35.1 µg/l), indicating a dilution effect from non-mine affected tributaries. Two dilution  
492 behaviours were observed in the main river, an abrupt reduction of Zn<sub>D</sub> after high  
493 concentrations were converged with relatively clean tributaries (from 3220.5µg/l (H7) and  
494 4252.3µg/l (H12) to 444.4µg/l (H10)), and a gradual decrease of Pb<sub>D</sub> (284.2µg/l (H12) to  
495 80.7µg/l (H10)) as dilution is likely to be related to the distribution of particulate matter from  
496 sediments<sup>70</sup>. Pb forms showed a higher fraction present as particulate (≥50%) in certain  
497 tributaries (e.g. H12, H11, H8, H3, H1) where dissolved organic carbon (H12 and H11>16  
498 mg/l) and other complexing compounds like bicarbonates (DIC in H8, H3 and H1>22 mg/l) or  
499 hydroxides (e.g. Fe(OH)<sub>3</sub> and Al(OH)<sub>3</sub>) may bind Pb. Thus, knowing solubility and speciation  
500 properties of Zn and Pb could help in the explanation of their mobility. Low solubility of Pb  
501 conceals high concentrations released at diffuse sources, due to binding to particulates<sup>71</sup>. In

502 addition, sorption properties also affect metal dynamics, for instance, Pb has a greater affinity  
503 for binding to dissolved organic matter and surface reactive mineral complexes, as reflected  
504 in higher fractions present as particulate forms. These mechanisms are fundamentally  
505 associated with metal speciation, bioavailability and toxicity. The bioavailability of Zn and Pb  
506 in Hebden Beck has been assessed, revealing quality standard failures of Pb and Zn  
507 throughout the catchment at all monitoring sites <sup>39</sup>.

508 Calculations of the dissolved Zn and Pb being transported downstream to the River Wharfe  
509 indicate annual loads of 0.2 tonnes/year of Pb and 2.9 tonnes/year of Zn. Although, these  
510 loads might increase depending on physical or chemical conditions caused by seasonal  
511 variations or particular flow conditions. Several studies in river systems have reported that  
512 metals associated with suspended sediments can make a major contribution to the total load  
513 of metals <sup>72, 73</sup>. Horowitz<sup>74</sup> compared and contrasted metal concentrations in suspended and  
514 bottom sediments versus dissolved levels, results indicated that bottom sediment  
515 concentrations were more than 100,000 (5 orders of magnitude) times higher than dissolved  
516 levels. Applying this approach in a downstream site (H1), higher results were observed for Pb  
517 (>430,000) and to a lesser extent for Zn (>17,000). Considering the importance of sediments  
518 in the transport and cycling of metals further work is needed in Hebden Beck. Comparisons  
519 between maximum measured concentrations of total dissolved metal concentrations in the  
520 main channel (site H2) with established regulatory limits for metals indicated that maximum  
521 annual concentrations of Al<sub>D</sub> (188 µg/l), Fe<sub>D</sub> (657 µg/l), Ba<sub>D</sub> (306 µg/l) and Sr<sub>D</sub> (356.4 µg/l)  
522 were within established regulatory limits (e.g. Al: 200 µg/l; Fe: 1000 µg/l; Ba: 1000 µg/l; Sr:  
523 1500 µg/l). Conversely, maximum annual concentrations of Pb<sub>D</sub> (87 µg/l) and Zn<sub>D</sub> (607 µg/l),  
524 when compared with environmental quality standards (EQS) showed maximum exceedances  
525 of 1200% with likely ecological effects <sup>75, 76</sup> (Table 4).

526 Metal toxicity and bioavailability are mainly controlled by metal concentrations, pH conditions  
527 and concentration of organic matter. Although the last two factors might have a stronger effect  
528 on biotic communities. For example, Ramsey<sup>77</sup> reported that soil acidity and organic matter  
529 concentration exerted stronger effects on plant and microbial community than metals. Thus,  
530 discriminating the influence of these key factors in biological processes is important from the  
531 perspective of dealing ecotoxicological effects of metals and potential restoration efforts <sup>5, 78</sup>.  
532 In this context, environmentally friendly and cost-effective techniques such as bioremediation  
533 have been developed for heavy metal removal/recovery where microbial remediation is  
534 particularly used in mine drainages due to the ability of microorganisms to generate alkalinity  
535 and immobilise metals <sup>79</sup>. For instance, in the UK the sulphur-reducing bacteria (e.g.  
536 *Desulfovibrio vulgaris*) has been used in the treatment of mine drainage due to its diverse

537 metabolic strategies to reduce sulphate ( $\text{SO}_4^{2-}$ ) to hydrogen sulphide ( $\text{H}_2\text{S}$ ), other elements  
538 like iron ( $\text{Fe(III)}$ ), oxygen and compounds like nitrate and nitrite and fumarate<sup>80</sup>. Aquatic  
539 organisms such as diatoms and invertebrates have an important role as biomonitors and  
540 bioindicators for assessing the impact of metal pollution. These organisms together with  
541 established EU-WFD classification tools and diversity indices (e.g. ASPT, N-TAXA) were used  
542 in a preliminary assessment of this catchment, however the effects of elevated metal levels  
543 were unclear<sup>81</sup>. Studies in neutral mine drainage carried out by Byrne et al.<sup>82</sup> indicated that  
544 the use of standard macroinvertebrate biotic and diversity indices (EU-WFD tools) could lead  
545 to erroneous classifications of aquatic ecosystem health. These results revealed that failure  
546 in the interpretation of biogeochemical interactions could lead to inaccurate analysis of  
547 organisms at risk of exposure, hence ineffective management decisions. Thus, the  
548 assessment of metal effects on living organisms is complex since biota might have different  
549 responses according to physiological processes, and because metals are subject to a range  
550 of factors affecting their level of reactivity, toxicity and bioavailability<sup>15</sup>. Considering WFD  
551 goals, more nuanced approaches are needed for assessing metals and their ecological  
552 effects. In this context, current chemistry data and updated chemical speciation tools will be  
553 used in future work to assess metal availability and toxicity and advising improvements to river  
554 basin management plans.

## 555 6. Conclusions

- 556 • Underlying limestone bedrock controls pH and weathering reactions, and therefore metal  
557 mobility within such catchments.
- 558 • Mobilisation of Zn and Pb have a strong dependence on site specific biogeochemistry and  
559 hydrological conditions. No dependence of Zn with flow suggests that  $\text{Zn}_D$  has no kinetic  
560 limitations on Zn mineral dissolution, whereas  $\text{Pb}_D$  varied according with flow variations,  
561 reflecting its tendency to be complexed with colloidal or particulate forms.
- 562 • Point sources are regular contributors of  $\text{Zn}_D$  despite flow fluctuations, while diffuse sources  
563 like spoil wastes produced higher contribution of  $\text{Zn}_D$  and  $\text{Pb}_D$  in overflow conditions after  
564 dry periods.
- 565 • Not all contributions of  $\text{Zn}_D$  and  $\text{Pb}_D$  are derived directly from oxidation of sphalerite and  
566 galena. Mineralogical and geochemical analysis revealed the contribution of secondary  
567 minerals such as smithsonite and cerussite, which are continuously leaching into the river  
568 and represent an added complexity for future remediation.
- 569 • Metal pollution results in a large area of catchment having a depleted chemical status with  
570 likely effects on the aquatic ecology.

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