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1	The impairment of river systems by metal mine contamination: A review
2	including remediation options
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24 Abstract

25 Contamination of aquatic environments as a consequence of deep metal 26 mining for Pb, Zn, Cu, Cd and Fe is of widespread international concern. 27 Pollution resulting from metal mining activities can result in significant 28 environmental and ecological degradation and can pose serious risks to 29 human health through contamination of food and drinking water. This paper 30 provides a review of the impacts of deep metal mine water discharges on 31 riverine sedimentology, hydrology and ecology and explores strategies for the 32 restoration of rivers draining historically abandoned metal mines. 33 34 Physical processes of mine waste dispersal are relatively well understood. 35 Chemical processes are more complex and much research is now focussed 36 on understanding geochemical and mineralogical controls on metal 37 attenuation and release. Recent advances in numerical modelling and 38 geochemical tracing techniques offer the possibility of identifying present and 39 predicted future patterns of contamination at the catchment scale. 40 41 The character of mine water has been extensively studied. However, 42 documented impacts on aquatic ecosystems can vary widely depending on a 43 range of hydroclimatological and geochemical factors. Numerous studies 44 have shown that the majority of the annual metal flux in rivers draining mining-45 impacted regions occurs during the summer and autumn months as a result of 46 water table drawdown, sulphide oxidation and dissolution and flushing of 47 metal salts during subsequent storm periods. There have been few highresolution studies of stormflow hydrochemistry, despite the importance of highflows in the translocation of mine wastes.

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51 A growing number of studies have documented chronic and acute toxic 52 effects of mine water contaminants, based on both field and laboratory 53 research, with specific reference to riverine macroinvertebrates. Common 54 bioindices have been used to examine the impacts of mine water 55 contaminants on macroinvertebrate ecology, although the success of these 56 indices has been mixed. Sublethal biomonitoring techniques, as distinct from 57 traditional laboratory bioassays with lethal endpoints, have gained 58 prominence as a means of detecting behavioural and physiological responses 59 of an organism to pulses of contaminants. The development of Biotic Ligand 60 Models (BLMs) has allowed organism physiology and important 61 environmental parameters to be factored into assessments of metal toxicity. 62 63 The strategies and technologies available for mine water remediation are 64 considered and key knowledge gaps are highlighted. Passive remediation 65 technologies offer a low cost and sustainable alternative to chemical 66 treatment of deep metal mine discharges. However, at present, these systems 67 generally fail to remove toxic metals associated with metal mine drainage to 68 an acceptable standard. New phytoremediation techniques offer the possibility 69 of immobilisation and extraction of toxic metals in mine spoil and 70 contaminated soils.

71

72 We conclude by identifying key recommendations for future research:

73 (1) Researchers and regulators should consider bioavailable metal fractions in 74 contaminated sediments, as opposed to total metal concentrations, if 75 sediment ecotoxicity is to be accurately measured. In addition, more 76 studies should make use of new spectroscopic techniques (e.g., XANES) 77 capable of providing more detailed information on metal speciation and, 78 therefore, sediment ecotoxicity. 79 (2) There is a need for better sampling and monitoring of toxic metal 80 concentrations and fluxes during stormflows in mining-impacted river 81 systems, especially given future predicted increases in stormflow 82 occurrence. In addition, further research is required to help understand the 83 potential toxicological impacts of stormflows in mining-impacted 84 catchments. 85 (3) Further research is required to develop biological indices to identify the 86 impacts of mine water contamination on macroinvertebrate communities. 87 (4) New substrates and techniques for remediation of metal-rich mine waters 88 are currently being investigated and pilot studies undertaken in the 89 laboratory and field. Many show promising results at the laboratory scale 90 but large-scale pilot treatment plants are required to test the efficiency and 91 long-term sustainability under field conditions. 92 (5) An interdisciplinary approach, incorporating the collaborative expertise and 93 knowledge regarding sedimentological / geological, hydrological, chemical 94 and ecological consequences of active and historic deep metal mining, is

95 advocated and should be utilised for effective river basin management and

96 the remediation and restoration of impacted sites.

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98	Keywords: metal mine, acid mine drainage, river sediment, flood
99	hydrochemistry, benthic macroinvertebrate, mine remediation
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122 **1. Introduction**

123 Environmental impacts of mining on aquatic ecosystems have received

- 124 increasing attention in recent years (Gray, 1998; Smolders et al., 2003; Olias
- 125 et al., 2004; Batty et al., 2010). Acidic drainage associated with the
- abandonment of coal mining activity has been a particular focus of research
- 127 (Banks and Banks, 2001). Contaminated discharge from abandoned metal
- mines and their spoil heaps has received less attention, reflecting the highly
- 129 variable responses associated with the complex and frequently site-specific
- 130 hydrogeological context of each, and the highly variable hydrogeochemical
- 131 characteristics of the discharge (Environment Agency, 2008a). However,
- 132 metal mine discharges have resulted in the severe degradation of many rivers
- across the globe (Gray, 1998; Gundersen and Stiennes, 2001; Olias *et al.*,
- 134 2004; Sola *et al.*, 2004; Poulton *et al.*, 2010).
- 135

136 Metal mining regions occur on all continents except Antarctica and even 137 extend to the continental shelf in certain areas where former floodplains have 138 been submerged by sea-level rise resulting from global warming (Aleva, 139 1985). As a consequence, significant contamination of the landscape, 140 including riverine and riparian habitats, has been reported internationally 141 (Smolders et al., 2003; Asta et al., 2007; Edraki et al., 2005; Gilchrist et al., 142 2009; Brumbaugh *et al.*, 2010). The most severely contaminated discharges 143 typically occur shortly after abandonment of a site, when artificial dewatering 144 has ceased and groundwater levels recover (Robb, 1994). Rising oxygenated 145 groundwater within deep mines interacts with metal sulphides in exposed 146 rockfaces, generating a leachate, typically characterised by low pH and high

147 concentrations of dissolved toxic metals and sulphates (Braungardt et al.,

148 2003; Gilchrist et al., 2009). Where the water table reaches the surface,

149 leachate may enter rivers and lakes as drainage from mine shafts and mine

150 drainage levels (adits), whilst rainwater may infiltrate through surface spoil

151 heaps and tailings to enter streams and other surface water bodies.

152

153 Within riverine systems receiving metalliferrous drainage, the composition and 154 health of plant and animal communities can be severely impaired through the 155 combined toxicity of reactive metals in both the water column and sediments, 156 sulphates and acidity (Sola et al., 2004; Schmitt et al., 2007; Batty et al., 157 2010; Chapa-Vargas et al., 2010). Aqueous metal concentrations generally 158 decline downstream of contaminated sources due to the precipitation of oxide, 159 hydroxide and sulphate phases, and co-precipitation or sorption of metals 160 onto these phases (Hudson-Edwards et al., 1999b). However, iron hydroxide 161 'ochre' and other metal precipitates can cover the entire river bed in extreme 162 instances and degrade habitat guality and important breeding and feeding 163 areas for instream organisms (Batty, 2005; Mayes et al., 2008). Chronic 164 contamination of riverine systems can be exacerbated by episodic flood 165 events (Bradley 1984; Hudson-Edwards et al. 1999a; Dennis et al. 2009) or 166 by the failure of tailing dams (Hudson-Edwards et al., 2003; Macklin et al., 167 2003; Sola et al., 2004). Such events have led to significant ecological 168 degradation in many regions of the world and have severely impacted 169 communities dependent on local rivers and their floodplains for food and 170 livelihood (Macklin et al., 2006; Taylor et al., 2010).

171

172 Environmental degradation resulting from metal mining is not restricted to 173 regions of the world where recent or active mineral exploitation is occurring. In 174 the UK, metal mining reached its peak in the mid-nineteenth century when, for 175 a time, the UK was the largest lead, tin and copper producer in the world 176 (Lewin and Macklin, 1987). Following a global reduction in metal prices 177 associated with the discovery of large deposits of lead and copper in the 178 Iberian Peninsula, South America and Australia during the late 19th and early 20th centuries, a decline of metal mining occurred throughout the UK. Today, 179 180 the number of abandoned metal mines in England and Wales is estimated at 181 over 3,000 (Jarvis et al., 2007). The historical legacy of these mines is still 182 present in the landscape in the form of spoil heaps, abandoned adits and 183 shafts, and derelict structures. The historical metal mining industry, long 184 forgotten and often far removed from manufacturing centre's, has left a 185 significant legacy of environmental contamination which will persist for 186 centuries to millennia (Environment Agency, 2002; Macklin et al., 2006). 187 Approximately 20% of all water quality objective failures in England and 188 Wales are due to drainage from abandoned metal mines (Environment 189 Agency, 2006). The severity of the problem is underscored by the view of the 190 Environment Agency of England and Wales that metal mine drainage poses 191 the most serious threat to water quality objectives after diffuse agricultural 192 pollution (Environment Agency, 2006).

193

Since the 1960s, concerns over the environmental impacts of historic metal
mining activities have gained increasing significance and this is reflected in
the growing body of literature on the topic (e.g., Macklin *et al.*, 2006; Batty *et*

197 al., 2010). However, due to the highly variable nature of environmental 198 degradation of surface waters draining metal mines and the site-specific 199 nature of many impacts, the literature is scattered through a wide range of 200 published sources (Wolkersdorfer, 2004). Unlike most review papers to date, 201 which largely focus on specific environmental compartments in relative 202 isolation to the wider aquatic ecosystem, this review paper aims to use an 203 interdisciplinary perspective to critically review: (1) the sedimentological, 204 hydrological and ecological impacts of metal mining activities; and (2) the 205 potential for remediation of metal mine sites and the existing remediation 206 technologies available.

207

208 The review is organised into 5 main sections. Mine water chemistry has been 209 studied extensively (e.g., Younger et al., 2002) and is generally well 210 understood. Therefore, the purpose of section 2 is to provide a brief overview 211 of the primary variables influencing the generation and character of metal 212 mine drainage. There have been several systematic reviews of the 213 sedimentological impacts of mining on the fluvial environment which have 214 documented the physical and chemical factors controlling metal dispersal and 215 storage in mining affected rivers systems (Lewin and Macklin, 1987; Macklin, 216 1996; Miller, 1997). In addition, new technologies and approaches to help 217 control and remediate sediment contamination have been widely considered 218 (e.g., Macklin et al., 2006). Section 3 provides a review of the recent 219 developments centred on new spectroscopic methods for the measurement of 220 metal mobility and speciation, and evaluate the performance of sediment environmental quality standards. Section 4 of this review examines the 221

222 catchment hydrological factors which influence the character of metal mine 223 drainage in fluvial systems and discusses the important role of stormflows in 224 transporting mine wastes from mine sites. In section 5, the ecological impacts 225 associated with metal mines are examined with specific reference to benthic 226 macroinvertebrate communities. While a significant body of research has 227 been devoted to examining impacts on fish communities (Hallare et al., 2010), 228 the benthic lifestyle of macroinvertebrates makes them more representative of 229 local environmental conditions, and, therefore, more reliable indicators of 230 biological stress. Previous reviews by Gerhardt (1993) and Batty et al. (2010) 231 have considered the impact of toxic metals and acidity on macroinvertebrates. 232 The present review builds on previous reviews by considering new 233 developments in biomonitoring techniques and sublethal measurements of 234 toxicity assessment. In the final section, remediation practices and 235 technologies to treat metal mine discharges are evaluated. In each of the four 236 key review sections (sedimentology, hydrology, ecology, remediation), we 237 highlight the key research gaps that remain and identify opportunities for 238 future research.

239

Given that previous reviews have considered the environmental impacts
associated with deep and surficial coal mining (Robb, 1994; Banks and
Banks, 2001; Younger, 2002), and in particular acid mine drainage (Robb and
Robinson, 1995; Banks *et al.*, 1997; Gray, 1997), this review focuses on the
impact of deep metal mines on riverine ecosystems with a particular emphasis
on the following widely exploited metals: lead (Pb), zinc (Zn), copper (Cu),
cadmium (Cd) and iron (Fe). All of these metals frequently occur at high

247 concentrations within waters draining metal mines (Novotny, 1995; Younger et 248 al., 2002). The review has broad geographical significance, but highlights 249 several case studies from the UK to illustrate some of the historic impacts of 250 metal mining activities on riverine ecosystems. Two search strategies were 251 used to identify relevant empirical papers. First, key word and title searches of 252 electronic databases were undertaken independently by the authors before 253 comparing results. The databases searched were: ASFA Aquatic Sciences, 254 Biological Sciences, Science Direct, SCOPUS, Toxline, Web of Science and 255 Zetoc. The key search words were: metal mine, heavy metals, toxic metals, 256 acid mine drainage, river sediment, flood hydrochemistry, benthic 257 macroinvertebrate, mine remediation and environmental quality. Databases 258 were searched from inception to December 2010. Second, relevant 259 references within any identified papers were followed up. Searches were 260 limited to papers published in English.

261

262 **2. Mine water chemistry**

263 Sulphidic minerals such as galena (lead sulphide - PbS), sphalerite (zinc 264 sulphide – ZnS) and pyrite (iron disulphide – FeS₂) are amongst the most 265 commonly mined metal ores (Novotny, 1995). These minerals are formed 266 under reducing conditions in the absence of oxygen and remain chemically 267 stable in dry, anoxic and high pressure environments deep underground. 268 However, these solid phases become chemically unstable when they are 269 exposed to the atmosphere (oxygen and water) through natural weathering 270 processes and long-term landform evolution or anthropogenic activities such as mining (Johnson, 2003). A series of complex biogeochemical reactions 271

272 occurs in sulphide weathering processes, leading to the generation of a 273 potentially toxic leachate and its release into the environment (Figure 1: 274 Younger et al., 2002; Johnson, 2003; Evangelou and Zhang, 1995). The 275 leachate generated during the sulphide weathering process is complex and is 276 often referred to as acid mine drainage (AMD) or acid rock drainage (ARD). It 277 is most commonly characterised by high levels of dissolved toxic metals and 278 sulphates and low pH (Robb and Robinson, 1995; Braungardt et al., 2003). 279 However, it should be noted that metal mine discharges are not always acidic 280 (Banks et al., 2002). In general, an increase in pyrite content of the country 281 rock results in greater acidity; an increase in base-metal sulphides results in 282 greater toxic metal concentrations; while an increase in carbonate and silicate 283 content can result in highly alkaline waters (Oyarzun et al., 2003; Alderton et 284 al., 2005; Cidu and Mereu, 2007). In the UK, much of central and north Wales 285 is underlain by Lower Palaeozoic shales and mudstones with low 286 concentrations of base materials (Evans and Adams, 1975). As a result, many 287 of the headwater streams of the region have low acid-buffering capability, 288 resulting in extremely acidic discharges containing high levels of dissolved 289 toxic metals (Abdullah and Royle, 1972; Fuge et al., 1991; Boult et al., 1994; 290 Neal et al., 2005). In contrast, in those parts of the English Peak District 291 where carbonate lithology predominates (Carboniferous Limestone), neutral to 292 basic mine discharges are common and these have significantly lower 293 concentrations of dissolved toxic metals (Smith et al., 2003). Aside from 294 lithology and mineralogy, the character of mine water pollution can vary 295 considerably between regions as a result of the grain size distribution of 296 tailings and spoil (Hawkins, 2004), the exposed mineral surface area

297 (Younger *et al.*, 2002), the concentration of reactants such as dissolved

298 oxygen (Wilkin, 2008), and microbial activity (Hallberg and Johnson, 2005;

299 Natarajan *et al.*, 2006; Balci, 2008). The highly variable nature of water

300 chemistry associated with metal mine discharges is outlined in **Table 1**.

301

302 3. Sedimentological impacts

303 During the lifetime of a deep metal mine, ore extraction and processing can 304 release vast quantities of solid waste into the riverine environment (Bird *et al.*, 305 2010). Even after mine abandonment, erosion of material from mine spoil and

306 tailings can continue to introduce contaminated solid wastes into river

307 channels and floodplains for many decades (Macklin et al., 2003; Walling et

308 *al.*, 2003; Miller *et al.*, 2004; Dennis *et al.*, 2009). These solid wastes can

309 have a significant impact on the geochemistry of channel and floodplain

310 sediments (e.g., Aleksander-Kwaterczak and Helios-Rybicka, 2009; Byrne et

311 *al.*, 2010) and also physical and chemical dispersion patterns of toxic metals

312 (e.g., Hudson-Edwards *et al.*, 1999b; Dennis *et al.*, 2009).

313

314 3.1 Sediment geochemistry

Gross contamination of fluvial sediments both within the channel and on the floodplain has been reported in most metal mining regions of the world (**Table 2**), with metal concentrations in sediments usually being several orders of magnitude greater than that in the water column (Macklin *et al.*, 2006). Metal concentrations are greatest in the fine sediment fraction and, in particular, in the clay-silt fraction (< 63 μ m; Lewin and Macklin, 1987; Foster and Charlesworth, 1996; Stone and Droppo, 1994; Dennis *et al.*, 2003; Förstner, 322 2004). This reflects the higher surface area per unit mass of smaller particles,
323 and the ion-exchange capacity of silt and clay-sized fractions (which include
324 clay minerals, iron hydroxides, manganese oxides, and organic matter in

325 various states of humification).

326

327 Metal speciation is essential to assess geochemical phases and the mobility 328 of potentially toxic elements in contaminated sediments (Tokalioglu et al., 329 2003). Until fairly recently, most investigations of sediment metal 330 concentrations have used strong chemicals (e.g., HNO₃, HCl, HClO₄, HF) to 331 extract the total amount of metals in the sediment, often leading to 332 oversimplified interpretations that do not take sediment complexity into 333 account (Linge, 2008). Metals in sediments exist in various geochemical 334 phases which reflect the degree to which they can be re-mobilised from the 335 sediment. For this reason, chemical sequential extraction procedures (SEPs) 336 capable of identifying contaminant partitioning have become increasingly 337 popular over total dissolution of the sediment achieved by single extractions 338 (e.g., Tessier et al. 1979; Rauret et al., 1999). Various extraction media have 339 been used to target specific geochemical phases, including electrolytes 340 (CaCl₂, MgCl₂), pH buffers of weak acids (acetic, oxalic acid), chelating 341 agents (EDTA, DTPA) and reducing agents (NH₂OH). In many metal mining 342 regions, the impact on sediment geochemistry has been to increase the 343 proportion of toxic metals in the more mobile (bioavailable) geochemical 344 phases. Studies have identified cadmium (Licheng and Guijiu, 1996; Morillo et 345 al., 2002; Vasile and Vladescu, 2010), copper (Jain, 2004), zinc (Morillo et al., 2002; Galan et al., 2003; Aleksander-Kwaterczak and Helios-Rybicka, 2009; 346

Naji *et al.*, 2010) and lead (Byrne *et al.*, 2010) to be highly elevated in the
acid-soluble phases. The largest proportion of metals is usually found in the
reducible phase bound to Fe and Mn oxides (Macklin and Dowsett, 1989).
Copper has been found to associate largely with organic matter in the
oxidisable phase (Licheng and Guijiu, 1996).

352

353 Sequential extraction procedures have allowed the chemical mobility and 354 toxicological risk posed by contaminated sediments to be established allowing 355 resource managers to prioritise areas for remediation. However, a number of 356 doubts concerning the accuracy of selective chemical extraction schemes 357 have been expressed (Linge, 2008). Particular concerns are whether the 358 chemical extractant may attack phases other than those expected; and 359 whether liberated metals may become associated with another sediment 360 phase rather than staying in solution (Burton, 2010). The multitude of 361 extraction techniques and media used can also lead to great variability in 362 results and, in some instances, limits the ability to make direct comparisons 363 between studies. Since the early 1990s, molecular scale techniques to study 364 elemental binding have become more popular. X-ray absorption spectroscopy 365 (XAS) techniques such as X-ray absorption near edge structure (XANES) and 366 extended X-ray absorption fine structure (EXAFS) have allowed analysis at 367 the molecular level and direct evidence of surface composition and bonding 368 characteristics of mining-derived sediments (Esbrí et al., 2010; Van Damme et 369 al., 2010). By investigating metal speciation at the atomic level, it is possible 370 to establish metal toxicity, mobility and bioavailability with far greater accuracy 371 than can be achieved using chemical extraction methods. This allows

372 scientists and environmental managers to more accurately gauge the impact373 of toxic metals on ecosystems and human health.

374

375 3.2 Physical dispersion and downstream attenuation processes

376 The influx of large volumes of contaminated material into river systems can

377 significantly alter local sediment transport and deposition and affect chemical

378 processes that operate at and beneath the river-bed surface (Gilbert, 1917;

Lewin *et al.*, 1977; Wood and Armitage, 1997). A number of reviews have

380 historically considered the hydrogeomorphic response of riverine systems to

mining activities (e.g., Miller, 1997; Macklin *et al.*, 2006) and as a result only

382 limited coverage is provided here. Based on research in the UK, Lewin and

Macklin (1987) suggested that disturbances of the river channel due to mining

384 can be categorized as involving processes of '*passive dispersal*' and '*active*

385 transformation'.

386

383

387 Passive Dispersal

388 During *passive dispersal*, mine waste is transported from the mine site with no 389 significant alteration of the prevailing sediment load of the river. Changes can 390 occur in depositional environments, with slow flowing and deep pools being 391 preferential sites for the deposition of contaminant-enriched fine sediment (< 392 2000 μ m). Transport of coarse sediment (> 2000 μ m) may be limited to 393 modest and high flow events. However, fine sediments may be transported 394 under a range of different flows, including extended periods of base-flow. In-395 channel sediment contamination generally decreases downstream from the 396 contaminant source at rates that vary between systems but which, in many

cases, are negatively exponential (Lewin and Macklin, 1987). This pattern is 397 398 functionally related to the hydraulic sorting of sediment based on density and 399 size of ore particles (e.g., galena is more dense than sphalerite and smaller 400 grains travel less fast than coarser grains - Wolfenden and Lewin 1978); 401 dilution by uncontaminated sediments (Marcus, 1987); hydrogeochemical 402 reactions (Hudson-Edwards et al., 1996); and biological uptake (Lewin and 403 Macklin, 1987). In many cases, good fits between metal concentration and 404 distance downstream of mining input can be achieved using regression 405 analysis (Wolfenden and Lewin, 1977; Lewin and Macklin, 1987) or non-linear 406 mixing models which incorporate clean and contaminated sediment sources 407 within a river catchment (Marcus, 1987). However, these models are often 408 specific to both the individual metal and the catchment for which they were 409 developed (Dawson and Macklin, 1998; Miller, 1997). Movement of sediment 410 can also occur in large-scale bed forms or 'slugs', which have been identified 411 as associated with highs and lows in an otherwise downward trending metal 412 concentration with distance downstream (Miller, 1997).

413

414 Active Transformation

Active transformation occurs in association with a significant increase in the
input of mining debris to the fluvial system. This may cause intrinsic
thresholds to be exceeded and, consequently, lead to a local loss of hydraulic
or geomorphological equilibrium that manifests itself in changes in channel
character (Lewin and Macklin, 1987). The type, rate and magnitude of
erosional and depositional processes can change (Miller, 1997). Channel
aggradation may be associated with sediment inputs from active mining and

422 channel degradation may occur after mining has ceased (Gilbert, 1917; 423 Knighton, 1991). Meandering channels may be transformed into braided 424 forms (Warburton et al., 2002). Other depositional features can include scroll 425 bars that arise from rapid accretion of sequentially developing point bars as a 426 response to high sediment loads and channel migration, and substantial 427 overbank floodplain deposits, particularly where overbank splays lead to 428 avulsion channels that cross the flood plain (Miller, 1997; Walling et al., 2003; 429 Dennis et al., 2009).

430

431 Toxic metal contaminants can be extremely persistent within the environment 432 and can remain stored within floodplain deposits for decades to millennia 433 (Miller, 1997). Since the 1970's, a significant amount of research has 434 focussed on the role of historical metal mining in the contamination of 435 floodplains (**Table 2**). Analyses of floodplain overbank sediments in the River 436 Ouse catchment in northeast England revealed contaminated sedimentary 437 successions reflecting over 2000 years of lead and zinc mining (Hudson-438 Edwards et al., 1999a). It has been estimated that over 55% of the 439 agriculturally important River Swale floodplain, a tributary of the Ouse, is 440 significantly contaminated by toxic metals (Brewer et al., 2005). It has been 441 estimated that approximately 28% of the lead produced in the Swale 442 catchment remains within channel and floodplain sediments. At present rates 443 of valley-bottom reworking through channel migration and erosion, it may take 444 in excess of 5,000 years for all of the metal-rich sediment to be exported from 445 the catchment (Dennis et al., 2009). These studies indicate that large areas of agricultural land are potentially contaminated and that there may be long-term 446

health concerns for those ingesting contaminants via crops produced on this
land (Albering *et al.*, 1999; Conesa *et al.*, 2010).

449

450 Recent advances in geochemical tracing techniques and numerical modelling 451 have led to improved understanding and predictability of dispersal rates and 452 patterns of sediment-associated toxic metal contamination. Owens et al. 453 (1999) used geochemical fingerprinting to identify the proportion of sediment 454 from mining areas in the River Ouse catchment, UK. Using isotope 455 signatures, several studies have differentiated specific geographical sources 456 in mining-affected catchments (Hudson-Edwards et al., 1999a; Bird et al., 457 2010). Bird et al. (2010) were able to discriminate between sediments derived 458 from mine waste and river sediments using lead isotope signatures. They 459 surmised that approximately 30% of the sediment load of the lower River 460 Danube was derived from mining. Numerical modelling techniques now allow 461 the prediction of contamination patterns in river catchments now and in the 462 future. For example, the catchment sediment model TRACER has been 463 applied to identify sediment contamination 'hot spots' in the River Swale 464 catchment, UK (Coulthard and Macklin, 2003). The model also revealed that 465 over 200 years after the cessation of mining activities, over 70% of the 466 deposited contaminants remain in the Swale catchment.

467

468 3.3 Chemical dispersion and attenuation processes

469 Chemical transportation processes in sediments of metal mining-affected

470 rivers become increasingly important after the closure and abandonment of

471 deep mines (Lewin and Macklin, 1987; Bradley et al., 1995). Toxic metals can

472 be attenuated downstream of a mining input through pH buffering, acid 473 neutralisation, and precipitation and adsorption reactions (Routh and 474 Ikramuddin, 1996; Ford et al., 1997; Lee et al., 2002; Ren and Packman, 475 2004). The often termed 'master variable' for determining metal speciation in 476 aquatic systems is pH (Kelly, 1988; Younger et al., 2002). As pH increases, 477 aqueous metal species generally display an increasing tendency to precipitate 478 as carbonate, oxide, hydroxide, phosphate, silicate or hydroxysulphate 479 minerals (Salomons, 1993). The effects of increasing pH below mine 480 discharges can be seen in some rivers by changes in precipitate mineralogy, 481 with proximal capture by iron hydroxides and distal capture by aluminium 482 oxides (e.g., Munk et al., 2002). Therefore, a major control on metal 483 attenuation, acid production and stream pH at abandoned mine sites is the 484 amount of carbonate minerals present in the surrounding geology. Carbonate 485 minerals such as calcite, dolomite and siderite weather quickly and can buffer 486 pH and act as adsorption sites for dissolved toxic metals. Non-carbonate 487 minerals weather slowly and, where they predominate, can be extremely slow 488 to react to changes in pH (Wilkin, 2008). The precipitation of solid-form metals 489 limits the concentration of metals which are transported through the aquatic 490 system as free ion species (Enid Martinez and McBride, 1998). These 491 secondary minerals can also act as sorbents for dissolved metals (Enid-492 Martinez and McBride, 1998; Asta et al., 2007; Wilkin, 2008). Adsorption of 493 metals usually increases at higher pH so that substantial changes in dissolved 494 metal concentrations can occur with small changes in pH, typically over 1 -495 1.5 pH units (Salomons, 1993). Aside from pH, several other water quality 496 parameters can influence metal speciation, including the concentration of the

metal, presence of ligands, redox conditions, salinity, hardness, and the
presence of other metals (Novotny, 2003). High levels of salinity, hardness
and organic matter content are known to increase metal attenuation by
providing binding sites for metal sorption (Salomons, 1980; Dojlido and
Taboryska, 1991; Achterberg *et al.*, 2003).

502

503 Under invariant environmental conditions, sediment geochemical phases are 504 stable, chemical attenuation of metals will proceed at regular rates and, thus, 505 metals remain immobile in river bed sediments (Morillo et al., 2002). However, 506 sediments are not a permanent sink for metals and they may be released into 507 the water column when suitable conditions for dissolution occur. Several 508 studies have reported the mobilisation of reduced sediment-bound metals to 509 the water column under oxidising conditions, for example, during floods and 510 dredging activities (Calmano et al., 1993; Petersen et al., 1997; Kuwabara et 511 al., 2000; Zoumis et al., 2001; Butler, 2009; Knott et al., 2009). In sediments 512 from Hamburg harbour, Calmano et al. (1993) observed oxidation episodes to 513 decrease pH in the suspended sediments from 7 to 3.4, leading to the 514 mobilisation of zinc and cadmium. Similarly, oxidation of anoxic sediments 515 from Mulde reservoir, Germany, resulted in the mobilisation of zinc and 516 cadmium and redistribution of toxic metals to more bioavailable geochemical 517 phases (Zoumis et al., 2001). Mullinger (2004) reported diffuse discharges of 518 metals from bed sediments accounted for up to 40% of zinc, cadmium and 519 copper entering surface waters of the Cwm Rheidol mine, Wales. Bioturbation 520 (Zoumis et al., 2001) and changes in pH (Hermann and Neumann-Mahlkau, 521 1985), dissolved organic carbon (Butler, 2009), ionic concentration (Dojlido

and Taboryska, 1991), and the concentration of complexing agents

523 (Fergusson, 1990; Morillo *et al.*, 2002) have also been reported to lead to the

524 release of 'stored' toxic metals into the wider environment.

525

526 The contamination risk posed by toxic metals stored in aquatic sediments of 527 former and current industrial centres (including metal mining regions), and the 528 potential for these toxic metals to contaminate areas beyond the source of 529 contamination, has prompted many national regulatory authorities to introduce 530 sediment environmental quality standards (SEQS) (e.g., Environment Agency, 531 2008b) based on total metal concentrations in the sediment. The practical 532 application of SEQS is made difficult by a number of factors relating to the 533 nature of heavy metal pollutants, including variation in natural background 534 concentrations, the existence of chemical species, the concentrations of 535 physico-chemical parameters, variations in organism sensitivity, and the fact 536 that some heavy metals are essential elements for organisms (Comber et al., 537 2008). In order to classify accurately the ecological status of rivers impacted 538 by metal mining, sediment assessments may need to be unique to each river 539 catchment and incorporate: background metal concentrations, an assessment 540 of bioavailable fractions, and concurrent water quality measurements 541 (including major ions) (Netzband et al. 2007; Brils 2008; Förstner 2009). As 542 far as is known by the authors, most national monitoring and assessment 543 programmes for freshwater systems measure total metal concentrations in 544 sediments rather than the concentration of metals in different geochemical 545 phases. Measurement of total quantities of metals in sediment provides little 546 information regarding their ecotoxicity and their potential mobility. With the

achievement of Good Ecological Status (GES) at the centre of many
environmental improvement programmes (e.g., to comply with the European
Water framework Directive), it is argued that measurement of bioavailable
metals in the sediment, which can interact relatively easily with aquatic
organisms, would provide a more comprehensive and robust assessment of
ecological risk. In this respect, there is a real risk that such programmes are
failing to meet their own objectives.

554

555 **4. Hydrological impacts**

556 The generation of mine water pollution is a product of many factors including 557 local mineralogy, lithology, contaminant source area, and biogeochemical 558 reactions (Younger et al., 2002). The character of mine water pollution in 559 surface waters is strongly influenced by a wide range of hydroclimatological 560 factors (including rainfall characteristics), land use (both catchment-wide and 561 any changes associated with spoil heaps), seasonality, antecedent conditions 562 to rainfall or snow-melt (particularly soil and spoil moisture content but also 563 temperature), dominant hydrological transport pathways, and stream 564 discharge (Gammons et al., 2005; Canovas et al., 2008). Once released to 565 the water column, metals can move through the aquatic environment, 566 resulting in impaired water quality in reaches of a river or estuary that were 567 unaffected directly by deep mine drainage. Released metals can also interact 568 with aquatic animals, resulting in the deterioration of aquatic ecosystem health 569 (Farag *et al.*, 1998).

570

571 Traditionally, discharge has been seen as a master variable driving river 572 hydrochemistry (Bradley and Lewin, 1982). Heavy metal ion concentrations in 573 rivers are generally thought to be greatest during low flows and lowest 574 coinciding with high flows, when uncontaminated runoff dilutes solute 575 concentrations (Webb and Walling, 1983). Since the 1970s, many 576 researchers have documented the effects of seasonal variability in stream 577 discharge on toxic metal concentrations (e.g., Grimshaw et al., 1976; Keith et 578 al., 2001; Sullivan and Drewer, 2001; Nagorski et al., 2003; Desbarats and 579 Dirom, 2005; Hammarstrom et al., 2005). Annual patterns (hysteresis 580 patterns) of dissolved metal concentrations are apparent in many rivers, 581 reflecting the flushing of oxidised sulphides accumulated over dry summer 582 (low flow) months (Canovas et al., 2008). Many researchers have noted 583 maximum toxic metal concentrations as occurring during the first heavy rains 584 of the hydrological year, during the autumn (Bradley and Lewin, 1982; Bird, 585 1987; Boult et al., 1994; Braungardt et al., 2003; Desbarats and Dirom, 2005; 586 Olias et al., 2004; Mighanetara et al., 2009). Contaminant concentrations 587 typically decrease in winter and increase gradually through spring and 588 summer as a result of increased sulphide oxidation and evaporation. 589 Therefore, the timing of maximum contaminant flux will be largely a function of 590 hydroclimatology, catchment characteristics and the minerals present at a 591 mine site. 592

593 It is understood that a major part of element transfer in rivers takes place

594 during short episodes of high river flow, i.e. floods (Sanden *et al.*, 1997).

595 However, to date, very little research has been directed towards detailing toxic

596 metal fluxes and hydrochemical variability during individual high flow events in 597 former mining regions (Grimshaw et al., 1976; Bradley and Lewin, 1982; 598 Sanden et al., 1997; Dawson and Macklin, 1998; Lambing et al., 1999; Wirt et 599 al., 1999; Keith et al., 2001; Gammons et al., 2005; Canovas et al., 2008). 600 One of the earliest studies by Grimshaw et al. (1976), on the River Ystwyth, 601 Wales, observed hysteretic behaviour in the relation between metal 602 concentrations and discharge, whereby metal concentrations increased on the 603 rising limb of the flood hydrograph and decreased on the falling limb. 604 associated with flushing and exhaustion (or dilution), respectively. This 605 general pattern has also been reported in a number of more recent studies 606 (e.g., Keith et al., 2001; Canovas et al., 2008; Byrne et al., 2009). In some 607 instances, the source of metals in the initial flush was metal sulphates 608 accumulated on the surface of mine waste (Keith et al., 2001) or 609 contaminated groundwater efflux from mine portals (Canovas et al., 2008). 610 Metal attenuation on the falling limb is principally due to rain-water dilution 611 and the fact that the available contaminant are scavenged in the first flush 612 (Canovas et al., 2008). The frequent occurrence of peak iron, manganese and 613 aluminium concentrations on the falling limb of the hydrograph indicates that 614 adsorption onto, or precipitation with, iron solids may be an important toxic 615 metal attenuation mechanism during stormflow events in some rivers (Lee et 616 al., 2002; Asta et al., 2007; Byrne et al., 2009).

617

618 The mobilisation and transport of mine wastes during stormflows and the

619 consequent contamination of agricultural lands is an important issue for

620 environmental managers of former metal mining regions (Dennis *et al.*, 2003;

621 Connelly, 2009). During the 1990s, there was a marked increased interest in 622 toxic metal contamination in floodplains in the UK following a number of 623 devastating floods and an increased focus on the potential effects of climate 624 change on hydrological regimes and sediment transport dynamics (Table 2). 625 The autumn and winter floods of 2000-2001 across a substantial part of 626 Europe caused large-scale remobilisation and deposition of contaminated 627 sediments in floodplains and farm-land (Dennis et al., 2003; Macklin et al., 628 2003: Macklin et al., 2006). In future, predicted increases in the frequency and 629 magnitude of floods as a function of climate change may result in increased 630 mobilisation and deposition of toxic metals in floodplains across Europe 631 (Macklin et al., 2006; Environment Agency, 2008b; Förstner and Salomons, 632 2008). Therefore, there is a need to monitor and assess stormflow events and 633 river hydrochemistry in detail in order to quantify metal fluxes with reasonable 634 levels of accuracy in order to allow environmental managers to prioritise areas 635 for remediation.

636

637 Aside from contamination of floodplains, the large-scale movement of mine 638 waste during stormflow events has significance for aquatic ecosystem health. 639 The highly elevated toxic metal concentrations during stormflows undoubtedly 640 cause harm to aquatic communities and degrade biological quality (Wolz et 641 al., 2009). The long-term effects of these transient conditions can be 642 established through investigations of aquatic ecosystem health. However, the 643 added or individual impact of stormflow events is still largely unknown due to 644 the difficulty of measuring it. Predicted increases in the frequency and 645 magnitude of floods across Europe due to climate change (Wilby et al., 2006)

have put an emphasis on bridging the knowledge gap between the physical
remobilisation of contaminants during stormflows and the potential
toxicological impacts (Wolz *et al.*, 2009). Understanding the toxicological
impacts of stormflows will be important in the achievement of environmental
quality standards in mining-affected river catchments.

651

652 Most metal mines are associated with significant volumes of waste material 653 deposited as surface spoil heaps and tailings. The hydrological behaviour of 654 these waste deposits can be significantly different to the wider catchment due 655 to the alteration of local surface and sub-surface flow pathways (Younger et 656 al., 2002). Considering the important role of spoil material in the production of 657 metal contaminants, comparatively little research has been undertaken into 658 flow pathways and contaminant generating processes within mine spoil. Due 659 to the artificial stratification and the discontinuities in permeability that occur 660 within spoil heaps, they often have 'perched aguifers' that lie well above the 661 underlying bedrock, producing unique flow paths (Younger et al., 2002). The 662 development of a water table in mine spoil depends on the predominant 663 lithology of the spoil. For example, sandstone generally forms highly 664 permeable spoil whereas mudstone produces spoil of low permeability. Highly 665 permeable spoil can contain as much as 25% or more of ore as fines or 666 solutes (Davies and Thornton, 1983). Where rainfall infiltration-excess is 667 typical, because, for instance, fine-grained material produces a surface seal, 668 surface runoff will be the predominant flow path (Younger et al., 2002). This 669 will, through gully erosion, transfer large quantities of contaminated solids into 670 the local water course.

671

672 Changes in flow paths and direction within mine spoil can occur slowly 673 through the seasons or more rapidly during rainfall events as different flow 674 paths become active with the fluctuation of perched water tables (Walling and 675 Webb, 1980). Differential hydrology can induce variability in toxic metal 676 speciation in mine spoils and tailings (Kovacs et al., 2006). Generally, 677 oxidation of sulphide minerals occurs in a shallow oxidation zone near the 678 surface of the spoil (Jurjovec et al., 2002). Dissolution and flushing of these 679 oxidised metals can then occur during wet periods (Navarro et al., 2008). 680 Several studies of metal flushing during storms have reported the importance 681 of weathered metal salts on and near the surface of mine spoil as responsible 682 for increasing metal concentrations during runoff (Canovas et al., 2008; Byrne 683 et al., 2009). Below the oxidation zone, a zone of transition from saturated to 684 unsaturated sediments typically occurs, often characterised by a 'hard pan' of 685 metal precipitates (Romero et al., 2007). Toxic metals can be attenuated in 686 the mine spoil through a series of precipitation, co-precipitation and 687 adsorption reactions. Reducing conditions in saturated sediments can lead to 688 the formation of insoluble metal sulphides. pH buffering can occur in the 689 shallow oxidizing zone with secondary-phase precipitation occurring near the 690 deeper saturated zone (McGregor et al., 1998). In order to effectively plan for 691 mine site remediation, it is essential that mine spoils and tailings are 692 characterised in terms of mineralogy, metal speciation and hydrology, 693 especially where contamination of groundwater is an issue. Such information 694 is necessary to understand the mechanisms controlling the release and 695 attenuation of metals at these sites.

696

697 5. Ecological impacts of metal mine contamination on macroinvertebrate 698 communities

As early as the 1960s, the adverse impacts of mining activities on

macroinvertebrates were being acknowledged (Reish and Gerlinger, 1964).

701 Metal mine drainage can severely impact aquatic ecosystems by affecting

primary and secondary production, nutrient cycling, energy flow and

decomposition (Stoertz et al., 2002; Knott et al., 2009; Younger and

704 Wolkersdorfer, 2004; Batty et al., 2010). Freshwater macroinvertebrates fulfil

important roles in the river ecosystem, being vital food sources for many

aquatic and terrestrial predators and playing a significant part in the cycling of

707 organic matter and nutrients (Gerhardt, 1993). The pivotal position of benthic

708 macroinvertebrates in aquatic food webs means that negative impacts on

them can have widespread consequences within aquatic and terrestrial food-

710 webs for primary producers, predators and the wider ecosystem. As a result,

711 macroinvertebrates have increasingly been used as indicators of stream

ecosystem health associated with metal mining (e.g., Batty *et al.*, 2010;

713 Poulton *et al.*, 2010).

714

5.1 Changes in community composition

A wide range of changes to macroinvertebrate community structure and composition have been reported in the scientific literature associated with metal mining activities. Reductions in abundance, number of taxa and biodiversity are common impacts reported in association with metal miningactivities internationally (e.g., Willis, 1985; Gray, 1998; Amisah and Cowx, 721 2000; Watanabe et al., 2000; Hirst et al., 2002; Kiffney and Clements, 2003) 722 (Table 3). Investigations have generally revealed that some 723 macroinvertebrate taxa display a tolerance or sensitivity to contamination 724 (Table 3). Whilst investigating contaminated stretches of two rivers in Ohio, 725 USA, Winner et al. (1980) hypothesised that habitats heavily polluted with 726 toxic metals may be dominated by Chironomidae (Diptera – true fly larvae); 727 moderately polluted habitats by Chironomidae and Trichoptera (caddisfly); 728 and minimally or unpolluted habitats by caddisflies and Ephemeroptera 729 (mayfly). Armitage et al. (1980; 2007) examined macroinvertebrate species 730 composition of the mining impacted River Nent. Diptera and Plecoptera 731 (stonefly) were the dominant orders observed in the river system. Trichoptera 732 and mayfly (Ephemeroptera) were not abundant and seemed particularly 733 sensitive to the mine water pollution. In contaminated reaches of the River 734 Vascao, Portugal, the number of predators increased and the number of EPT 735 taxa (Ephemeroptera – Plecoptera - Trichoptera) decreased, probably 736 reflecting the presence of thick layers of metal hydroxides on the river 737 substrate (Gerhardt et al., 2004). Sites subject to severe AMD contamination 738 showed high levels of biodiversity due to high species richness of the tolerant 739 species. In general, the order of toxicity of metal mine contamination to the 740 most common macroinvertebrate orders is: Ephemeroptera > Trichoptera > 741 Plecoptera > Diptera. However, there can be considerable variability in metal 742 tolerance between macroinvertebrate taxa and species. For example, 743 Ephemeroptera are generally considered to be highly sensitive to metal 744 contamination despite some species (e.g., Baetis rhodani and Caenis cf. 745 *luctuosa*) being reported to display some tolerance to metal contaminants

747 Gerhardt et al., 2005b). Several authors have reported impacts of mine water 748 contamination on ecosystem function (**Table 3**), including reduced secondary 749 production (Carlisle and Clements, 2005; Woodcock and Huryn, 2007), and a 750 reduction in leaf matter (detritus) decomposition rates and microbial 751 respiration (Kiffney and Clements, 2003; Carlisle and Clements, 2005). 752 753 Relatively predictable changes in macroinvertebrate community structure as a 754 result of pollution (e.g., decreased abundance and biodiversity, elimination of 755 sensitive taxa) have led to the development of a number of biotic and diversity 756 indices (e.g., Shannon, 1948; Berger and Parker, 1970). However, the 757 performance of biological indices / metrics appear to vary widely when applied 758 to mine water contaminated sites (Smolders et al., 2003; Van Damme et al., 759 2008; Chadwick and Canton, 1984; Willis, 1985; Chadwick et al., 1986; Rhea 760 et al., 2006). Variability in success is likely to be a function of the complicated 761 interplay between the mine water components, other water quality 762 parameters, and natural tolerances and sensitivities of organisms. Gray and 763 Delaney (2008) suggest a modification of the Acid Waters Indicator 764 Community (AWIC) index (Davy-Bowker et al., 2005) to incorporate metal 765 toxicity may be required. However, such a revision would also need to 766 address the pH bias in the calibration data and the (possibly) inaccurate 767 grouping of macroinvertebrates in sensitivity groups. A revision of the 768 Biological Monitoring Working Party (BMWP) system (Biological Monitoring 769 Working Party, 1978), based on species' tolerance to acidity and metal contamination, has also been suggested (Gray and Delaney, 2008) and some

(Roline, 1988; Beltman et al., 1999; Gower et al., 1994; Gerhardt et al., 2004;

746

770

success has been achieved using a multi-metric approach by considering
multiple biological metrics simultaneously (e.g., Clews and Ormerod, 2009).
Clearly, there is scope for a biological index designed specifically for detecting
the impacts of mine water contamination on aquatic communities. However,
such an index would need to incorporate the effects on a community of
multiple environmental stressors, the most important of which are probably
dissolved metals and acidity.

778

5.2 Changes in macroinvertebrate physiology and behaviour

780 More subtle community alterations as a result of physiological or behavioural 781 changes are less easy to diagnose (Younger and Wolkersdorfer, 2004) (Table 782 3). For example, Petersen and Petersen (1983) reported anomalies in the 783 construction of filter feeding nets of Hydropsychidae (Trichoptera) in rivers 784 affected by a gradient of toxic metal pollution. Disruption of silk-spinning by 785 contamination caused the caddisfly to spend more time in open habitats 786 repairing the structure and thus more vulnerable to potential predators. Vuori 787 (1994) observed metal exposure to affect the territorial behaviour of 788 Hydropsychidae, relaxing levels of interspecific competition and increasing 789 susceptibility to predation. Brinkman and Johnston (2008) reported decreased 790 moulting rates (*Rhithrogena hageni*: Ephemeroptera) after exposure to high 791 levels of copper, cadmium and zinc. In an experimental stream study, 792 Clements et al. (1989) reported that high copper doses increased predation 793 pressure, so much that the numbers of caddisfly, mayfly and chironomids 794 were dramatically reduced. Maltby and Naylor (1990) found high zinc 795 concentrations significantly impacted *Gammarus pulex* reproduction by 796 causing a reduction in energy absorption and an increase in the number of

797 broods aborted. Other behavioural responses reported associated with metal 798 mine contamination include increased drift rates, physical avoidance of 799 contaminated sediments, reduced burrowing / burial rates (Leland et al., 1989; 800 Roper et al., 1995) and reduced leaf litter processing rates and microbial 801 respiration (Kiffney and Clements, 2003; Carlisle and Clements, 2005). Many 802 of the species specific differences reported within the literature have been 803 attributed to trophic status with herbivores and detritivores typically being 804 more sensitive to contamination than predators (Leland *et al.*, 1989; 805 Schultheis et al., 1997; Gerhardt et al., 2004; Poulton et al., 2010). Acute 806 metal contamination can induce deformities and mutations of head and 807 feeding structure in macroinvertebrate fauna (e.g., Groenendijk et al., 1998; 808 Vermeulen et al., 2000; Groenendijk et al., 2002; De Bisthoven et al., 2005). 809 Both zinc and lead have been implicated as teratogens (inducing deformities 810 as a result of chronic exposure during the lifetime of the organism) and as a 811 mutagen (inducing deformities in offspring due to DNA damage in parents 812 from chronic exposure) in Chironomus riparius (Chironomidae) (Martinez et 813 al., 2004).

814

More recent studies have made use of biomonitoring techniques which are
capable of detecting sublethal behavioural and physiological responses in an
organism when exposed to a contaminant (e.g., De Bisthoven *et al.*, 2004;
Gerhardt *et al.*, 2004; Gerhardt *et al.*, 2005a; De Bisthoven *et al.*, 2006;
Gerhardt, 2007; Macedo-Sousa *et al.*, 2007) (**Table 3**). A conceptual
Stepwise Stress Model (SSM), proposed by Gerhardt *et al.* (2005a),
postulates that an organism will display a time-dependent sequence of

822 different regulatory and behavioural responses during exposure to

823 contaminants over a certain threshold. Several species have been found to

show a pH-dependent response to AMD involving, first, an increase in

locomotion, followed by an increase in ventilation (e.g., Gerhardt *et al.*, 2005a;

B26 De Bisthoven *et al.*, 2006). An increased ventilation rate reflects changes in

the organism's respiratory and physiological system, and may be due to

828 damage to gill membranes or nerve tissues. Locomotory activity probably

829 represents an avoidance strategy from potentially toxic conditions.

830 Importantly, biomonitoring methods integrate biochemical and physiological

processes and so are a more comprehensive method than single biochemical

or physiological parameters. In combination with the Stepwise Stress Model,

833 online biomonitoring offers the possibility of a graduated 'early warning'

system for the detection of pollution waves (Gerhardt *et al.*, 2005a).

835

836 5.3 Metal bioaccumulation in macroinvertebrates

837 A significant body of research has concentrated on evaluating the 838 bioaccumulation of toxic metals in macroinvertebrates as a measure of the 839 bioavailability of contaminants (e.g., Farag et al., 1998; Smolders et al., 2003; 840 Yi et al., 2008). Metals which are bioaccumulated by organisms and plants 841 can be concentrated or magnified in the food chain (Sola et al., 2004) (Table 842 3). Benthic primary producers and decomposers are known to accumulate 843 significant amounts of metals with little or no deleterious effects (Farag et al., 844 1998; Sanchez et al., 1998). These metals can be transferred to herbivorous 845 and detritivorous macroinvertebrates which in turn can transfer the metals to higher trophic levels (Younger and Wolkersdorfer, 2004). Metal accumulation 846

847 can vary between species, depending on a great number of physiological (e.g. 848 cuticle type, the presence or absence of external plate gills, the processes 849 which control metal distribution in the cell) and behavioural factors such as an 850 organisms feeding strategy, contact with benthic sediments, larval stage and 851 size (Dressing et al., 1982; Farag et al., 1998; Goodyear and McNeill, 1999; 852 Sola and Prat, 2006; Cid et al., 2010). Metal intake can take place through 853 direct exposure to metals in surface and pore waters or indirectly via food 854 supply. Those metals which, through their chemistry, are almost completely 855 sediment-bound (Fe, Mn, Pb, Al), will usually be most important for particle 856 feeders. Metal intake in the tissue takes place at a cell membrane, typically in 857 the gill or gut, depending on whether the metal is in solution in the 858 surrounding water body or if it was ingested with food. A range of 859 environmental factors determine the potential for metal bioaccumulation 860 including metal concentration in the surrounding water, water hardness, 861 presence of organic matter, feeding group and the ionic state of the metal 862 (Gower and Darlington, 1990; Farag et al., 1998; Sola and Prat, 2006). The 863 accumulation of metals in different organisms can also vary greatly as a result 864 of natural or evolved tolerance mechanisms (Spehar et al., 1978; Gower and 865 Darlington, 1990; Bahrndorff et al., 2006). For example, Plectrocnemia 866 conspersa (Trichoptera), common in streams in south-west England affected 867 by metal mine drainage were found to be tolerant of copper pollution (Gower 868 and Darlington 1990). Some controlled microcosm experiments have reported 869 tolerance to metal polluted sediments by Chironomus februarius 870 (Chironomidae) (Bahrndorff et al. 2006). Mechanisms of tolerance might be 871 methylation, increased metal excretion or decreased metallothionein
872 production. Metallothionein is a metal-binding protein with the principal 873 function of accumulating essential metals for normal metabolic processes 874 (Howard, 1998). Its presence facilitates the accumulation of toxic metals, 875 however decreased production of this protein may allow certain organisms to 876 accumulate lower amounts of toxic metals. Despite the great range of factors 877 which can affect metal bioaccumulation in organisms, bioaccumulation factors 878 (BAFs) which consider tissue metal concentration in relation to the 879 surrounding abiotic medium, are possibly a more robust biodiagnostic method 880 than measurement of metal concentrations in the water column and benthic 881 sediments. If water quality guidelines are to continue to be used, then 882 additional research will need to be undertaken to determine appropriate 883 guidelines (possibly above existing guidelines) to support aguatic 884 communities. In the future, metal bioaccumulation will need to be studied in a 885 greater range of macroinvertebrates in order to fully understand metal-886 organism interactions in aquatic systems. A review of metal bioaccumulation 887 studies by Goodyear and McNeill (1999) found that most studies primarily 888 considered Ephemeropteran and Dipteran taxa and especially collector-889 gatherer and predatory functional feeding groups / traits.

890

5.4 Effects of environmental parameters on the toxicity of mine discharges
Changes in some environmental parameters can affect the chemistry and,
therefore, the toxicity of metals to organisms. The effects of salinity, water

894 hardness and alkalinity on metal toxicity have been studied extensively (Stiff,

1971; Brkovic-Popovic and Popovic, 1977; Gauss *et al.*, 1985; Gower *et al.*,

896 1994; Yim *et al.*, 2006, Riba *et al.*, 2010 – **Table 3**). All of these studies

897 reported metal toxicity increases for macroinvertebrate and fish species under 898 low salinity, alkalinity and water hardness conditions. Increased metal toxicity 899 has also been reported at low turbidity (Garcia-Garcia and Nandini, 2006) and 900 DOM (dissolved organic material) levels (Gower et al., 1994). In river 901 systems, carbonate minerals, clay minerals and DOM act as sorption sites for 902 toxic metals and, therefore, high levels of these parameters help to reduce the 903 concentration of dissolved toxic metals in bioavailable forms. However, 904 bottom-dwelling organisms will take up sediment-bound metals through 905 ingestion.

906

907 While bioassay and microcosm studies have revealed much information on 908 metal ecotoxicity, a possible criticism of them could be that they are too 909 simplistic in seeking to evaluate the response of macroinvertebrate species or 910 communities to a single metal contaminant. In reality, most contaminated 911 mine waters will contain mixtures of different metals in solution (Table 3). The 912 simplest solution has been to assume the toxic effects of the metals present in 913 the mixture are additive (Vermeulen, 1995). However, the interaction between 914 metals can result in synergistic effects. For example, Hickey and Golding 915 (2002) reported total abundance of heptageniid mayflies, community 916 respiration and macroinvertebrate drift were most sensitive to solutions with a 917 mixture of zinc and copper. Clements (2004), in stream mesocosms, found 918 negative responses were generally greatest with zinc alone or with zinc and 919 cadmium. A possible explanation for this synergism is the physiological 920 inhibition of metal excretion by one of the metals, allowing the other metal(s) 921 to have greater toxic effects (Berninger and Pennanen, 1995). Mixtures of

922 metals have also been shown to have antagonistic effects. Morley et al. 923 (2002) found zinc and cadmium to have an antagonistic effect leading to 924 increased survival of the cercarial stage of the parasitic fluke *Diplostomum* 925 spathaceum. In some cases, antagonistic effects of metal mixtures are 926 probably related to competition between metal ions for common sites of 927 uptake (Younger and Wolkersdorfer, 2004). A study by Vermeulen (1995) 928 illustrated the difficulty in predicting how metal mixtures will affect metal 929 toxicity to organisms. Out of the 26 studies analysed, thirteen reported 930 synergistic effects, six reported antagonistic effects, and seven reported 931 additive effects. The problem of metal mixture toxicity is further compounded 932 by other water quality parameters such as hardness, salinity and organic 933 matter content. These parameters can increase or decrease metal toxicity and 934 comparable mixtures of metals can also show contrasting toxicity effects 935 between different groups, species and populations of organisms (Younger 936 and Wolkersdorfer, 2004).

937

938 The task of evaluating metal toxicity is made even more difficult when acidity 939 is considered. Most commonly, a decrease in pH will increase the amount of 940 toxic free metal ions due to changes in metal speciation, mobility and 941 bioavailability (Campbell and Stokes, 1985). However, at low pH, metals tend 942 to desorb from organisms due to competition with hydrogen ions for binding 943 sites (Gerhardt, 1993). The effects of low pH on stream biota in the absence 944 of dissolved metals can be lethal or sublethal, inducing a range of 945 physiological changes including an upset of the ionic balance across 946 organism membranes and hydrolysing of cellular components (Kelly, 1988).

Campbell and Stokes (1985) suggested acidity can affect metal-organism 947 948 interactions in two key ways. First, if a decrease in pH causes little change in 949 metal speciation and there is only weak binding of metals at biological 950 surfaces, the decrease in pH will decrease the toxicity of the metal due to 951 competition with hydrogen ions for binding sites. Second, if a decrease in pH 952 causes changes in speciation and there is strong binding at biological 953 surfaces, then acidification will increase metal availability and toxicity. In the 954 first instance, acidity will be the primary threat to ecosystems. In the second 955 scenario, low pH and high dissolved metals may both influence toxicity.

956

957 The multi-factor nature of contaminated mine discharges (acidity, dissolved 958 metals, metal precipitates, sulphates) and the natural variability in water 959 chemistry between regions means that metal toxicity can be highly variable. 960 Historically, ambient water quality criteria have specified permissible total or 961 dissolved metal concentrations even though metal toxicity is heavily 962 dependent on water chemistry (e.g., hardness, pH, DOM). The Biotic Ligand 963 Model (BLM) (Di Toro et al., 2001) was developed to predict metal toxicity by 964 incorporating basic principles of physiology and toxicology, and the effects of 965 water chemistry on metal speciation and bioavailability. The model has gained 966 widespread use amongst the scientific / academic and water industry 967 communities due to its potential for identifying water guality criteria and in 968 facilitating risk assessment of aquatic environments (Paguin et al., 2002). In 969 order to gain wider applicability and relevance, BLMs will need to be applied 970 to a wider range of organisms and pollutants in the future, and to be able to 971 incorporate metal mixtures into toxicity predictions (Niyogi and Wood, 2004).

972

973 6. Remediation of mining-impacted river systems

974 The prevention of contaminated discharge from mine sites is now required by 975 law in many countries (Macklin *et al.*, 2006). In the USA, the Clean Water Act

976 (1972) was established to minimise the impact of anthropogenic pressures

977 (including mining) on surface waters. In Europe, the adoption of the Water

978 Framework Directive (2000/60/EC), and subsequent Mining Waste Directive

979 (2006/21/EC), has necessitated the development of inventories of

980 contaminant impacts at active and abandoned mine sites (Hering *et al.*, 2010).

New legislation, based on a greater understanding of water quality and

982 ecological integrity issues arising from mine discharges, have prompted

983 research into remediation technologies aimed at reducing the environmental

984 impact of metal mines (PIRAMID Consortium, 2003).

985

986 Mine water remediation technologies can be broadly categorised into active 987 and passive treatment. Active treatment technologies are well established and 988 involve the utilisation of electrical energy and mechanised procedures (Jarvis 989 et al., 2006) and are dependent on continuous monitoring and maintenance 990 (Robb and Robinson, 1995). Traditional active treatment processes involve a 991 sequence of oxidation by physical or chemical means, the addition of alkaline 992 chemicals to raise pH and accelerate oxidation and precipitation of metals 993 (Robb and Robinson, 1995; Lund and McCullough, 2009), and settlement and 994 filtration (PIRAMID Consortium, 2003). However, active treatment incurs 995 substantial set-up, material and maintenance costs (PIRAMID Consortium, 996 2003). In response, passive remediation utilising natural physical, chemical

997 and biological processes and materials has found increasing favour over the 998 past 30 years (Geroni et al., 2009). Passive remediation systems use 999 naturally available energy (e.g., topographical gradient, metabolic energy, 1000 photosynthesis) to drive he remediative processes and have the principal 1001 advantages over active remediation of reduced set up and maintenance costs 1002 (Pulles and Heath, 2009). Some passive systems (e.g., wetlands) require 1003 significantly greater land area than active treatment systems; although they do 1004 not require costly reagents and incur less operational maintenance (Norton, 1005 1992; Hedin et al., 1994). Detailed characterisation of contaminant loading 1006 over a sufficiently long time period is required prior to implementation of 1007 treatment systems, including measurements of seasonal variation and the 1008 impact of episodic contaminant flushing events, e.g., associated with spate 1009 flows (Younger et al., 2005; Byrne et al., 2009). Equally important is the 1010 linking of all mine water sources with a treatment system. Many abandoned 1011 mine sites have substantial diffuse sources (Pirrie et al., 2003; Mayes et al., 1012 2008; Mighanetara et al., 2009; Byrne et al., 2010), including mine spoil and 1013 mobile metal fractions in the river bed. As a result it may be difficult to collect 1014 and route contaminated runoff to treatment areas.

1015

1016 Mine water treatment technologies have been extensively reviewed elsewhere

1017 (e.g., Brown *et al.*, 2002; Younger *et al.*, 2002; PIRAMID Consortium, 2003;

1018 Lottermoser, 2007) and so a brief overview is provided (Table 4). Both

1019 wetlands and Reducing and Alkalinity Producing Systems (RAPS) are now

1020 well established remediation technologies throughout North America (e.g.,

1021 Hedin *et al.*, 1994) and Europe (e.g., Whitehead and Prior, 2005) as passive

1022 treatment options for sulphate and Fe-rich, net-alkaline and net-acidic coal 1023 mine discharges (Batty and Younger, 2004). In anoxic systems, removal of 1024 toxic metals (e.g., zinc, lead, copper, cadmium) is hypothesised to occur 1025 through the formation of insoluble metal sulphides and carbonates (Younger 1026 et al., 2002 – See **Table 4**). In aerobic systems, some toxic metals can be 1027 removed either by direct precipitation as oxides and hydroxides or carbonate 1028 phases or by co-precipitation with iron, manganese and aluminium 1029 hydroxides. However, rates of toxic metal removal in these systems 1030 (particularly zinc) have, in general, proved insufficient in circum-neutral and 1031 net-alkaline mine waters, where chalcophile metals are the principal 1032 contaminants (Robb and Robinson, 1995; Nuttall and Younger, 2000). Some 1033 success has been achieved using variations of conventional calcite and 1034 organic-based treatment systems in laboratory-scale experiments (Nuttall and 1035 Younger, 2000; Rotting et al., 2007; Mayes et al., 2009). A large number of 1036 researchers have also demonstrated the potential for organic and inorganic 1037 sorbent media to remove toxic metals (Cui et al., 2006; Perkins et al., 2006; 1038 Madzivire et al., 2009; Mayes et al., 2009; Rieuwerts et al., 2009; Koukouzas 1039 et al., 2010; Vinod et al., 2010). However, many of these technologies are still 1040 at the experimental stage and will require further refinement and large-scale 1041 field pilot studies before their full potential is realised. Frequent blocking of 1042 filtering media with metal precipitates and rapid consumption of reactive 1043 surfaces limit the metal removal efficiency of many of these systems to very 1044 short time scales – hours to days in some instances (Younger et al., 2002). 1045

1046 Even with mine water treatment, the legacy of contamination in river 1047 sediments and floodplains will represent a significant secondary diffuse 1048 source of pollution long after other water quality parameters have improved to 1049 acceptable levels. Therefore, contaminated sediments of mining-affected 1050 rivers will continue to pose a serious threat to ecological integrity and the 1051 achievement of Good Chemical Status (GCS) and Good Ecological Status 1052 (GES) under the EU Water Framework Directive. The historical, preferred 1053 method of dealing with contaminated sediment is removal by dredging (Navar 1054 et al., 2004). This is an expensive and destructive process which may 1055 mobilise vast reservoirs of bioavailable metals as part of the process (Navar 1056 et al., 2004; Knott et al., 2009). Furthermore, the sediment removed still 1057 requires treatment and safe disposal. Recently, geochemical engineering 1058 approaches involving in-situ and ex-situ biological and chemical treatment of 1059 contaminated soils and sediments have gained attention as alternatives 1060 (Förstner, 2004), and some success has been achieved in the stabilisation and removal of toxic metals (Guangwei et al., 2009; Luoping et al., 2009; 1061 1062 Scanferla et al., 2009). However, the principal necessity for the protection of 1063 sediment and aquatic systems is considered to be the development of 1064 quidelines concerning sediment quality (Burton, 2010; Byrne et al., 2010). 1065 1066 Some efforts have focussed on the prevention of the generation of 1067 contaminated mine water, so-called source control techniques. Conventional 1068 techniques have focussed on physical and chemical stabilisation (Mendez

1069 and Maier, 2008). Physical stabilisation involves covering mine waste with

1070 inert material (e.g., clay, gravel) to reduce oxygen inflow and water ingress

1071 into the contaminated material (Gandy and Younger, 2003; Waygood and 1072 Ferriera, 2009). However, clay caps in arid and semi-arid regions have tended 1073 to crack from wetting and drying cycles resulting in the failure of the air-tight 1074 cap (Newson and Fahey, 2003). Chemical stabilisation is achieved by adding 1075 a resinous adhesive to form a crust over the mine waste, however, these also 1076 are prone to cracking and failure (Tordoff *et al.*, 2000). More recently, 1077 phytoremediation (phytoextraction and phytostabilisation) techniques have 1078 developed as less costly alternatives (Margues et al., 2009). 1079 Phytostabilisation creates a vegetative cap on the mine waste which 1080 immobilises metals by adsorption and accumulation in the rhizosphere 1081 (Mendez and Maier, 2008). Some success has been achieved in laboratory 1082 trials investigating reforestation of mine tailings using endemic tree species 1083 (Pollmann et al., 2009). Phytoextraction offers the possibility of recovery of 1084 metals through the hyperaccumulation of metals in plant tissues (Ernst, 2005). 1085 However, the long-term performance of these new strategies needs to be 1086 evaluated, as does the bioavailability of metals to wildlife which may feed on 1087 the vegetative covers.

1088

1089 **7. Synthesis and conclusions**

1090 This paper provides a critical synthesis of scientific literature related to the 1091 sedimentological, hydrological and ecological impacts of metal mining on 1092 aquatic ecosystems. It has also highlighted the potential for remediation of 1093 mine sites and provided an overview of current research and technological 1094 developments in this area.

1095

1096 The important role of sediments in the dispersal, storage and recycling of 1097 metal contaminants within the fluvial environment has been highlighted. 1098 Significant quantities of contaminated sediment are eroded and transported 1099 into aquatic systems from abandoned metal mines and both physical and 1100 chemical processes influence the distribution of toxic metals within riverine 1101 ecosystems. Physical dispersal processes are generally well understood and 1102 can be classified as passive or active (Lewin and Macklin, 1987), the latter 1103 prevailing when the addition of mine wastes to a river system results in a 1104 threshold crossing event and the collapse of geomorphological equilibrium. 1105 Under these circumstances, significant contamination of floodplains by toxic 1106 metals can occur, with long-term potential consequences for the environment, 1107 society and human health. However, recent advances in geochemical tracing 1108 techniques and numerical modelling have led to improved understanding and 1109 predictability of dispersal rates and patterns of sediment-associated toxic 1110 metal contamination (Coulthard and Macklin, 2003). Chemical dispersal of 1111 mine wastes tends to predominate after mine closure and four principal 1112 processes result in toxic metal attenuation downstream of inputs - pH 1113 buffering, acid neutralisation, precipitation and adsorption. However, river 1114 sediments are not a permanent store for toxic metals and they may be 1115 released into the water column if there are fluctuations in some important 1116 environmental parameters (i.e. pH and redox potential). As a result, 1117 establishing metal speciation, bioavailability and potential mobility is essential 1118 in order to prioritise sites for remediation. Recently, molecular scale 1119 techniques to study elemental binding have become more accessible to 1120 researchers. A greater number of geochemical studies should make use of

1121 these techniques to provide more accurate information on bonding 1122 characteristics of metals in sediments. Environmental regulators are 1123 beginning to acknowledge the central role of sediments in maintaining 1124 ecological guality in river systems. We have argued that the measurement / 1125 quantification of total metal concentrations, as is practiced by many 1126 regulators, provides limited information on the potential toxicity of sediments. 1127 Measurement of the bioavailable metal fraction within benthic sediments is 1128 considered a more accurate gauge of potential metal toxicity.

1129

1130 The character of metal mine drainage after it enters surface waters is affected 1131 by many factors including stream discharge, rainfall characteristics, conditions 1132 antecedent to rainfall-runoff events and season, and the interaction of a large 1133 permutation of processes which must be understood and quantified in order to 1134 mitigate effectively. Seasonal variability in metal concentrations is linked to 1135 oxidation and dissolution of metal sulphates, leading to elevated metal 1136 concentrations in summer and autumn months. At many mine sites, the 1137 transport of significant amounts of mine waste is limited to stormflows. 1138 Typically, hysteresis is evident in the relationship between metal 1139 concentrations and discharge. Peak metal concentrations are achieved before 1140 peak discharge, associated with the dissolution of surface oxidised material. 1141 Despite the importance of stormflows for the transport of mine wastes, little 1142 research has concentrated on investigating toxic metal fluxes and 1143 hydrochemical variability under these conditions. Predicted increases in the 1144 frequency and magnitude of floods as a function of climate change may result 1145 in increased mobilisation and deposition of toxic metals in floodplains across

1146 Europe. Stormflow hydrochemistry in rivers draining mine sites should be 1147 studied in more detail in order to quantify metal fluxes more accurately and 1148 allow environmental managers to prioritise areas for remediation. Toxic metal 1149 flushing during stormflows potentially impacts stream ecosystems by 1150 significantly increasing the toxicity of the river water, even if only for short time 1151 periods. More research is needed to help understand the potential 1152 toxicological impacts of stormflows in mining-affected river catchments. 1153 Relatively few studies have investigated mine spoil hydrology and metal 1154 attenuation and release processes. Environmental investigations at 1155 abandoned metal mine sites should include assessments of mine spoil in 1156 terms of mineralogy, metal speciation and hydrology, especially where 1157 contamination of groundwater is an issue.

1158

1159 Metal mine contaminants in river systems can have a variety of negative 1160 impacts on macroinvertebrate ecology and biology, including changes to community structure, physiological and behavioural impacts as well as direct 1161 1162 mortality. Typically, rivers heavily impacted by metal mine drainage have 1163 reduced species diversity and abundance, and tend to be dominated by 1164 Dipteran species. The order of toxicity in mining-impacted streams generally 1165 proceeds in the order Ephemeroptera > Trichoptera > Plecoptera > Diptera. 1166 Bioindices are used widely to guantify contaminant impacts on 1167 macroinvertebrate communities. However, there effectiveness in discerning 1168 the impacts of metal mine contamination is guestionable, with widely varying 1169 performance reported in the literature. The problem appears to be related to 1170 the multi-factor nature of mine discharges. Further research is required to

1171 develop a biological index specifically for the detection of the impacts of mine 1172 water contamination on macroinvertebrate communities and the wider 1173 ecosystem. Traditionally, laboratory bioassay experiments have been used to 1174 investigate metal and AMD toxicity, with organism mortality being the test 1175 endpoint. Recently, biomonitoring techniques capable of detecting sublethal 1176 behavioural and physiological responses in an organism have become 1177 popular (e.g., Gerhardt et al., 2004). They have the principal advantage over 1178 bioassays of integrating both biochemical and physiological processes. A 1179 major criticism of bioassay and microcosm studies is that they generally do 1180 not consider metal mixtures or the influence of other environmental 1181 parameters on metal toxicity. The development of the Biotic Ligand Model has 1182 allowed organism physiology and important environmental parameters to be 1183 factored into assessments of metal toxicity (Di Toro et al., 2001). However, to 1184 reach their full potential, BLMs will need to be applied to a wider range of 1185 organisms and pollutants, and they will need to be able to incorporate metal 1186 mixtures into toxicity predictions.

1187

1188 An increasing range of remediation technologies have been developed for the 1189 treatment of contaminated mine water which can be applied in a variety of 1190 topographical settings. Chemical treatment of mine waters is expensive and 1191 unsustainable over the substantial time periods treatment will be required. 1192 Passive remediation technologies offer a low cost and sustainable alternative. 1193 Passive systems for the treatment of coal mine discharges, where iron, 1194 sulphates and acidity are the principal contaminants, are considered proven 1195 technology. However, these systems generally fail to remove toxic metals

1196 (e.g., Zn, Pb, Cd), associated with metal mine discharges, to an acceptable 1197 standard. New substrates and techniques aimed at removing high 1198 concentrations of these toxic metals are being trialled and many show 1199 promise at the laboratory scale. However, large-scale pilot treatment plants 1200 are needed in order to develop these new systems and to test them in field-1201 relevant conditions. Even with mine water treatment, mine spoil and 1202 contaminated soils in mining regions will continue to pose a threat to water 1203 and ecological quality for many years into the future. New bio-based source 1204 control techniques such as phytoremediation offer the possibility of stabilising, 1205 immobilising and extracting toxic metals from soils at low cost, by using plants 1206 which hyper-accumulate toxic metals in their tissue. However, the long-term 1207 functioning and ecological impact of these new strategies needs to be 1208 evaluated.

1209

1210 A management approach which can draw on the expertise of separate but 1211 related and relevant disciplines such as hydrology, hydrochemistry, sediment 1212 geochemistry, fluvial geomorphology and aguatic ecology affords the 1213 opportunity for a more complete understanding of processes and impacts in 1214 mining-impacted river catchments. It is hoped that this review will help to 1215 contribute to our knowledge and understanding of the impacts of metal mining 1216 on aquatic ecosystems and highlight the usefulness of approaching such 1217 problems from a multi-disciplinary geographical point of view. 1218

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2213 Figure 1 The process of pyrite weathering in a deep metal mine. Four general 2214 equations describe the chemistry of pyrite weathering and the production of 2215 AMD - (1) The oxidation of pyrite by oxygen and water in atmospheric 2216 conditions to produce dissolved ferrous iron and sulphuric acid; (2) the 2217 oxidation of dissolved ferrous iron to ferric iron; (3) the hydrolysis of ferric iron 2218 with water to produce iron hydroxide precipitate (ochre) and acidity; (4) the 2219 oxidation of additional pyrite by the ferric iron generated in reaction (2) to 2220 produce dissolved ferrous iron and sulphuric acid. The acidic conditions 2221 generated during these processes can dissolve oxidised trace metals. The 2222 process is accelerated by the presence of sulphide and iron-oxidising 2223 bacteria.

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2225 **Table 1** A comparison of dissolved metal (mg/l), sulphate (mg/l) and pH concentrations from waters impacted by historical deep

2226 metal mining.

Location			Sample type	Pb	Zn	Cu	Cd	Fe	SO ₄	рН	Author(s)
Europe											
River Carnon, England			Mine drainage	<0.01 - 0.02	0.12 - 23	0.02 - 1.3	<0.01 - 0.02	<0.01 - 49	77 - 789	3.3 - 7.7	Neal et al. (2005)
River Tamar, England			Adit drainage	<0.01 - 0.17	<0.1 - 2.5	<0.01 - 1.4	<0.01 - 0.01	0.05 - 2.6	10 - 89	3.4 - 7.8	Mighanetara et al. (2009)
Funtana	Raminosa	Mining	Tailings	<0.01	0.08 - 34	<0.01 - 0.04	<0.01 - 0.85	0.02 - 0.25	22 - 1680	7.1 - 7.8	Cidu and Mereu (2007)
District, Italy			drainage								
Buchim	Mining	district,	Mine stream	0.03*	0.03*	0.62*	<0.01*	0.3*	-	5.1*	Alderton et al. (2005)
Macedonia											
Zletovo	Mining	District,	Adit drainage	0.06*	21.57*	0.46*	0.14*	98.2*	-	3.4*	Alderton et al. (2005)
Macedonia											
River Zletovska, Macedonia		Channel	<0.03 - 0.8	0.04 - 70.07	<0.01 - 1.05	<0.01 - 0.24	0.1 - 103.3	-	3.4 - 7.6	Alderton et al. (2005)	
River Bjorgasen, Norway		Channel	-	5.4 ^b	2.7 ^b	0.01 ^b	-	-	3.2 ^b	Gundersen and Stiennes (2001)	
Rio Tinto, Spain		Channel	0.1 - 2.4	0.3 - 420	0.05 - 240	-	-	2800 - 16000	1.4 - 7.6	Hudson-Edwards et al. (1999b)	
Troya Mine, Spain		Tailings pond	0.02 - 0.05	4.99 - 18.95	<0.01 - 0.03	0.01 - 0.03	0.04 - 0.33	-	-	Marques et al. (2001)	
River Odiel, Spain		Channel	<0.01 - 1.18	0.17 - 130.23	0.01 - 37.62	<0.01 - 0.38	0.03 - 262.71	50.7 - 3960	2.5 - 6.3	Olias et al. (2004)	
Tintillo River, Spain		Mine drainage	0.01 - 0.07	7.3 - 216	3.5 - 115	<0.01 - 0.51	264 - 1973	1300 - 11580	2.3 - 2.8	Sanchez Espana et al. (2006)	
Tinto Santa Rosa Mine, Spain		Mine drainage	<0.01 - 0.08	56 - 85	15 - 23	0.09 - 0.15	234 - 881	2704 - 4026	2.6 - 3.4	Asta et al. (2007)	
Fluminese Mining District, Spain	Mine water	<0.01 - 0.05	0.88 - 40	-	<0.01 - 0.09	<0.01 - 12	17 - 640	6.3 - 8.2	Cidu et al. (2007)		
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River Tawe, Wales	Channel	<0.01 - 0.15	0.01 - 8.8	<0.01 - 0.04	<0.01 - 0.16	-	-	-	Vivian and Massie (1977)		
River Rheidol, Wales	Channel	<0.01	0.08 - 0.29	-	<0.01	-	5.3 - 7.1	5.5 - 6.4	Fuge et al. (1991)		
River Yswyth, Wales	Channel	0.06 - 0.09	0.17 - 0.36	-	<0.01	-	nd - 5.3	4.1 – 4.6	Fuge et al. (1991)		
Cwm Rheidol Mine, Wales	Adit drainage	0.02 - 0.04	38 - 72	0.03 - 0.07	0.04 - 0.11	-	441 - 846	2.8 - 3.0	Fuge et al. (1991)		
Cwm Ystwyth Mine, Wales	Spoil drainage	0.29 - 3.3	1.5 - 4.6	<0.01	<0.01	-	nd	4.1*	Fuge et al. (1991)		
Cae Coch Pyrite Mine, Wales	Mine water	-	-	-	-	2261 ^b	6590 ^b	2.4 ^b	McGinness and Johnson (1993)		
River Goch, Wales	Channel	-	<0.01 - 4.19	<0.01 - 5.99	-	<0.01 - 25.98	-	2.3 - 7.7	Boult et al. (1994)		
Cwm Rheidol Mine, Wales	Spoil drainage	-	577 - 978	1.2 - 9.35	-	-	-	2.6 - 2.7	Johnson (2003)		
North America											
West Squaw Creek, USA	Channel	-	0.01 - 156	<0.01- 190	-	0.03 - 500	2.6 - 5100	2.4 - 6.9	Filipek et al. (1987)		
Richmond Mine, USA	Mine water	1 - 120	$0.06 - 23.5^{a}$	0.21 - 4.76ª	4 - 2110	2.47 – 79.7ª	14 - 760 ^a	-3.6 - 1.5	Nordstrom et al. (2000)		
Peru Creek, USA	Channel	-	0.55 - 1.89	0.05 - 0.22	-	0.08 - 0.5	29.6 - 73	4.7 - 5.9	Sullivan and Drever (2001)		
Boulder Creek, USA	Channel	<0.032*	0.469*	0.246*	<0.01*	2.82*	97.4*	3.3*	Keith et al. (2001)		
Black Foot River, USA	Channel	-	<0.2 - 535	<0.8 - 4	<0.5 - 2.6	<5 - 37	5.5 - 88.8	7.3 - 8.8	Nagorski et al. (2002)		
Phillips Mine, USA	Channel	<0.01	<0.01 - 0.17	0.02 - 3.13	-	0.16 - 42.4	25 - 368	2.3 – 6.5	Gilchrist et al. (2009)		
Australasia											
River Dee, Australia	Channel	<0.01 - 0.6	<0.01 - 10.4	<0.01 - 45.03	-	<0.01 - 74	340 - 5950	2.7 - 7.0	Edraki et al. (2005)		

Mt. Morgan Mine, Australia	Open pit	1.51*	21.97*	44.54*	-	253*	13600*	2.7*	Edraki et al. (2005)	
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nd = not detectable. * single observation. a grams per litre. b mean value

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2240 **Table 2** Comparison of metal concentrations (mg/kg) in channel and floodplain sediments from historic deep metal mining impacted

2241 rivers.

River location	Geomorphic-type	Grain size	Metal phase	Pb	Zn	Cu	Cd	Author(s)
	site	fraction	extracted					
Europe								
Red River, England	Channel	<2000 μm	Total	nd - 120	nd - 630	nd - 1320	-	Yim (1981)
River Derwent, England	Channel	<1000 µm	Total	96 - 3120	82 - 2760	-	0.6 - 13.8	Burrows and Whitton (1983)
River Derwent, England	Floodplain	<2000 µm	Total	131 - 1179	<10 - 1696	2.9 - 64	0.08 - 12.5	Bradley and Cox (1990)
River Tyne, England	Floodplain	<2000 µm	Total	615 - 2340	722 - 2340	11 - 42.5	2.6 - 8	Macklin et al. (1992)
River Swale, England	Floodplain	<63 µm	Total	56 - 5507	15 - 3066	-	1 - 18	Macklin et al. (1994)
River Allen, England	Channel	<170 µm	Total	2330*	1410*	-	-	Goodyear et al. (1996)
River Severn, England	Floodplain	<2000 μm	Total	23 - 204	173 - 936	30 - 67	0.35 - 6.4	Taylor (1996)
River Tees, England	Channel	<2000 μm	Total	522 - 6880	404 - 1920	20 - 77	0.95 - 5.95	Hudson-Edwards et al. (1997)
River Aire, England	Channel	<63 µm	Total	90 - 237	274 - 580	118 - 198	-	Walling et al. (2003)
River Swale, England	Floodplain	<63 µm	Total	10000*	14000*	-	7500*	Dennis et al. (2003)
River Calder, England	Channel	<63 µm	Total	199 - 343	397 - 907	141 - 235	-	Walling et al. (2003)
River Wear, England	Channel	<150 µm	Total	20 - 15000	40 - 1500	<10 - 340	-	Lord and Morgan (2003)
Dale Beck, England	Channel	<2000 μm	Total	13693*	442*	206*	-	Geer (2004)
River Avoca, Ireland	Channel	<1000 µm	Total	-	1520ª	674ª	-	Herr and Gray (1996)
River Mala Panew, Poland	Channel	<63 µm	Total	36 - 3309	126 - 11153	3.97 - 483	0.18 - 559	Aleksander-Kwaterczak and Helios-

								Rybicka (2009)
River Somes, Romania	Channel	<2000 µm	Total	28 - 6800	64 - 19600	12 - 8400	0.8 - 110	Macklin et al. (2003)
River Viseu, Romania	Floodplain	<2000 µm	total	17 - 850	110 - 2760	32 - 1000	0.5 - 17	Macklin et al. (2005)
Gezala Creek, Spain	Channel	<177 μm	Total	10.6 - 37630	216 - 25676	2.7 - 1691	0.22 - 45	Marques et al. (2001)
River Tinto, Spain	Channel	<2000 µm	Total	3200 - 16500	600 - 67300	1800 - 26500	<1 - 23	Galan et al. (2003)
River Odiel, Spain	Channel	<2000 µm	Total	1900 - 16600	1000 - 74600	3500 - 20900	1.4 – 10.2	Galan et al. (2003)
River Rheidol, Wales	Floodplain	<2000 µm	Total	291 - 2098	242 - 630	21 - 85	0.08 - 3.5	Davies and Lewin (1974)
River Tawe, Wales	Channel	<2000 µm	Total	63 - 6993	20 - 31199	34 - 2000	2 - 335	Vivian and Massie (1977)
River Rheidol, Wales	Floodplain	<210 µm	Total	813 - 1717	201 - 383	33 - 120	-	Wolfenden and Lewin (1977)
River Towy, Wales	Channel	<2000 µm	Total	36 - 5732	106 - 3722	44 - 259	0.78 - 83	Wolfenden and Lewin (1978)
River Twymyn, Wales	Channel	<2000 µm	Total	593 - 6411	159 - 6955	44 - 2557	1.5 - 44	Wolfenden and Lewin (1978)
River Ystywth, Wales	Floodplain	<2000 µm	Total	73 - 4646	123 - 1543	-	-	Lewin et al. (1983)
River Twymyn, Wales	Channel	<63 µm	Non-residual	1.1 - 2914	0.7 - 148	0.3 - 30	<0.01 - 0.9	Byrne et al. (2010)
North America								
West Squaw Creek, USA	Channel	<177 μm	Total	-	32 - 5940	254 - 4090	-	Filipek et al. (1987)
Black Foot River, USA	Channel	<63 µm	Total	1100 - 8700	1700 - 9600	1400 - 9900	<1 - 115	Nagorski et al. (2002)
River Cedar, USA	Channel	-	Total	4.5 - 420	9.75 - 2050	2.3 - 107	0.07 - 3.8	Ouyang et al. (2002)
Copper Mine Brook, USA	Channel	<1000 µm	Total	9.9 - 30	9 - 67	31 - 398	-	Gilchrist et al. (2009)
Australasia								
River Kangjiaxi, China	Channel	-	Non-residual	1154 - 8034	124 - 2319	23 - 209	2.6 - 41	Licheng and Guiju (1996)

nd = not detectable. * maximum value. ^b mean value.

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2255 **Table 3** Impacts of metal mine drainage on instream macroinvertebrates reported within the scientific literature. Types of studies

2256 are - ^a stream survey, ^b microcosm experiment and ^c laboratory bioassay.

Primary impact reported	Additional information	Author(s)
Community composition		
Shift in community structure	Clean sites dominated by Ephemeroptera and Plecoptera; moderately contaminated sites dominated by	Armitage (1980) ^a
	Plecoptera and Diptera; and heavily contaminated sites dominated by Diptera	
	Clean sites dominated by Ephemeroptera; moderately contaminated sites by Tricoptera; and heavily	Winner <i>et al</i> . (1980) ^a
	contaminated sites dominated by Diptera	
	Contaminated sites dominated by Orthocladiinae (Chironomidae) and species of net-spinning Tricoptera	Clements <i>et al</i> . (1992) ^a
	Contaminated sites dominated by Chironomidae	Gray (1998)ª
	Ephemeroptera reduced by > 75% in moderately contaminated streams	Clements et al. (2000) ^a
	Clean sites dominated by Stenopsychidae (Trichoptera); contaminated sites dominated by Chironomidae and	Watanabe et al. (2000) ^a
	Epeorus latifolium (Ephemeroptera)	
	Contaminated sites dominated by Chironomidae, Tubificidae, Baetidae and Simulidae	Marques <i>et al</i> .(2003) ^a
	Heavily contaminated sites dominated by Chironomidae	Smolders et al. (2003) ^a
	Dominance of predators in very acidic mining sites	Gerhardt <i>et al</i> . (2004) ^a
	Heavily contaminated sites characterised by high proportion of Chironominae and predatory Tanypodinae	Janssens de Bisthoven <i>et al</i> . (2005) ^a
Decrease in abundance	Reduction in abundance recorded	Willis (1985) ^a , Gray (1998) ^a , Hirst <i>et al</i> .
		(2002) ^a
	Ephemeroptera comprised less than 5% of individuals at one location	Clements <i>et al.</i> (1992) ^a

	Abundance significantly lower in experiments with metal mixtures and high predation pressure	Kiffney (1996) ^b
	Abundance positively related to stream alkalinity and pH	Malmqvist and Hoffsten (1999) ^a
	Ephemeroptera and Plecoptera particularly affected	Clements (2004) ^b
Decrease in number of taxa	Reduced number of taxa recorded	Willis (1985)ª, Kiffney (1996) ^b , Gray
		(1998) ^a
	Decrease most pronounced in low flow conditions	Clements <i>et al.</i> (1992) ^a
Decrease in EPT taxa	EPT richness positively related to stream pH	Malmqvist and Hoffsten (1999) ^a
	Near extinction of mayfly species	Hickey and Golding (2002) ^a
	Reduced number of EPT taxa recorded	Gerhardt <i>et al</i> . (2004) ^a
Decrease in species diversity	Reduced species diversity recorded	Amisah and Cowx (2000) ^a , Hirst et al.
		(2002) ^a
	Dominance of Chironomidae	Smolders et al. (2003) ^a
	Dominance of Chironomidae, Baetidae and Simulidae	Van Damme <i>et al</i> . (2008) ^a
Impaired ecosystem function	Microbial colonisation of leaf material and leaf decomposition inhibited by high Cd concentrations	Giesy <i>et al.</i> (1978) ^b
	Microbial activity and leaf decomposition rates significantly lower at contaminated sites	Carpenter et al. (1983) ^a
	Secondary production of shredders negatively associated with metal contamination; leaf decomposition rates	Carlisle and Clements (2005) ^a
	decreased; microbial respiration decreased	
	Reduced secondary production and organic matter storage	Woodcock and Huryn (2007) ^a
	Greater vulnerability of net-spinning Tricoptera to predation possibly due to spending more time in the open	Clements et al. (1989) ^b
	repairing capture nets	

Macroinvertebrate physiology

Physiological response	Differences in metal sensitivity related to trophic status; herbivores and detritivores more sensitive than	Leland <i>et al</i> . (1989)ª
	predators	
	Decrease in reproduction rates of Gammuras pulex (Gammaridae)	Maltby and Naylor (1990)°
	Differences in sensitivity related to trophic status; reduced leaf decomposition rates suggests shredders	Schultheis <i>et al</i> . (1997) ^a
	sensitive to pollution	
	Increase incident of deformity (mentum structure) in Chironomous riparius (Chironomidae)	Groenendijk <i>et al</i> . (1998)ª
	Increased incident of deformity (mentum structure) and decreased moulting success in Chironomous riparius	Vermeulen <i>et al</i> . (2000) ^c
	(Chironomidae)	
	pH-dependent decrease in locomotion of Atyaephyra desmaersti (Crustacea) in AMD solutions	Gerhardt <i>et al</i> . (2004)°
	Locomotion and ventilation of Choroterpes picteti (Leptophlebiidae) greater in acid only solutions than in AMD	Gerhardt <i>et al</i> . (2005a)°
	solutions	
	pH-dependent increase in locomotion and ventilation of Gambusia holbrooki (Crustacea) in AMD solutions	Gerhardt <i>et al</i> . (2005b) ^c
	pH-dependent decrease in locomotion and ventilation of Atyaephyra desmaresti (Crustacea) in AMD solutions	Janssens De Bisthoven <i>et al</i> . (2006)°
	Contaminated water causes higher locomotory activity in Lumbriculus variegates (Oligochaeta) than	Gerhardt (2007)°
	contaminated sediment	
	Decrease in pH and increase in dissolved metals caused decrease in locomotion and inhibition of feeding rate	Macedo-Sousa <i>et al</i> . (2007)°
	in Echinogammarus meridionalis (Crustacea)	
	Pulse of AMD caused early warning responses in Echinogammarus meridionalis (Crustacea) consisting of	Macedo-Sousa <i>et al</i> . (2008)°
	increased locomotion and subsequent increase in ventilitation	
	Average daily moulting rate of Rithrogena hageni (Heptageniidae) decreased after exposure to aqueous	Brinkman and Johnston (2008)°
	copper, cadmium and zinc	
Behavioural response	Anomalies in capture nets of Hydropsychidae	Petersen and Petersen (1983) ^a

	Decrease in burrowing rates and increase in crawling and drifting rates of Macomona liliana (Bivalve)	Roper <i>et al</i> . (1995)°
Morphological deformities	Cross-breeding of Chironomous riparius (Chironomidae) from contaminated and clean rivers revealed some	Groenendijk <i>et al</i> . (2002) ^b
	level of genetic adaptation to metals in offspring	
	Macroinvertebrate drift and respiration significant correlated with metal concentrations	Clements (2004) ^b
	Increased incident of adult and larval deformities in Chironomous tentans (Chironomidae)	Martinez <i>et al</i> . (2004)°
	Decreased locomotory activity of Chironomous sp. (Chironomidae) in AMD solutions	Janssens De Bisthoven <i>et al.</i> (2004)°
Metal bioaccumulation	Younger instars had higher metal concentrations than older instars	Krantzberg (1989)°
	Concentration of metals in Ephemeropteran species decreased in consecutive larval stages	Jop (1991)°
	Metal bioaccumulation dependent on feeding group; shredders and scrapers accumulated the highest metal	Farag <i>et al</i> . (1998)ª
	concentrations (biofilm contained more metals than sediments)	
	Whole-body metal concentrations of Hydropsyche sp. (Hydropsychidae) greater in species exposed to	DeNicola and Stapleton (2002) ^b
	dissolved metals than in species exposed to AMD precipitates	
	Chironomus februarius (Chironomidae) exhibited adaptation to and tolerance of metal-polluted sediments	Bahrndorff <i>et al.</i> (2006) ^b
	Macroinvertebrate metrics significantly correlated with metals in biofilm, suggesting biofilm is a better index	Rhea <i>et al.</i> (2006) ^a
	than macroinvertebrates for monitoring metal impacts on aquatic systems	
	Whole-body metal concentrations of Hydropsyche sp. (Hydropsychidae) were strongly positively correlated	Sola and Prat (2006) ^a
	with metal concentrations in water and sediment	
Effects of environmental parameters on the	ne toxicity of metal mine discharges	
Water hardness and alkalinity	Increased water hardness and alkalinity reduces metal toxicity in Chironomous tentans (Chironomidae)	Gauss <i>et al</i> . (1985) ^c
	Increasing water hardness reduces community sensitivity to metal contamination	Gower <i>et al</i> . (1994) ^a
	Increased water hardness reduces metal toxicity in Daphnia magna (Dapniidae)	Yim <i>et al.</i> (2006) ^c

	Metal mixtures	Abundance of heptageniidae, community respiration and macroinvertebrate drift were more sensitive to metal	Hickey and Golding (2002)°
		mixtures than single metal solutions	
		Survival of Diplostomum spathaceum (Diplostomatidae) greater in metal mixtures than in single metal solutions	Morley <i>et al.</i> (2002) ^c
		Community sensitivity greatest in combined metal mixtures compared to single metal solutions	Clements (2004) ^b
	Other parameters	Increased turbidity reduces metal toxicity to Cladocera by decreasing bioavailability of metals	Garcia-Garcia and Nandini (2006)°
		Inverse correlation between salinity and lesion index of gills in Ruditapes philippinarum (Bivalvia)	Riba <i>et al</i> . (2010) ^c
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2268 **Table 4** Typology of common passive mine water treatment units and source control techniques: indicating the nature of mine

2269 water drainage and the principal advantages and limitations of each method.

Name	Mine water type	Brief description	Advantages	Limitations	Example reference(s)					
Passive mine water treatment technologies										
Aerobic wetlands	Net alkaline	A system of shallow ponds,	Efficient Fe and Al removal;	Not suitable for highly toxic,	Robb and Robinson (1995);					
	ferruginous	cascades and vegetated	low maintenance	sulphate-rich and acidic mine	Johnson and Hallberg (2005)					
		substrate encourage aeration	requirement; cost-effective;	waters; large land surface						
		of mine waters and oxidation,	easy integration into	area requirement; occasional						
		hydrolysis and precipitation of	landscape and connection	removal of substrate						
		some heavy metals (mainly	with existing ecosystems	precipitates required						
		Fe and AI)								
Anaerobic wetlands	Net acidic ferruginous	A thick anoxic substrate of	Often used to neutralise	Not suitable for high toxic	Younger <i>et al</i> . (2002);					
	with high sulphate	saturated organic material	acidity and generate alkalinity	metal concentrations	Johnson and Hallberg (2005)					
	concentrations	neutralises acidity and	prior to discharge to aerobic	(especially Zn and Cd); large						
		generates alkalinity through	wetlands; efficient Fe and	land surface area						
		processes of bacterial	sulphate removal; some toxic	requirement; occasional						

		sulphate reduction and calcite	metals are removed through	removal of substrate	
		dissolution; heavy metals	precipitation of sulphides and	precipitates required; requires	
		(mainly Fe and Al) are	adsorption to organic matter;	high sulphate (>100 mg/l)	
		removed as precipitates	low maintenance	concentrations; often produce	
			requirement; cost-effective;	hydrogen sulphide gas	
			easy integration into		
			landscape and connection		
			with existing ecosystems		
Anoxic Limestone	Net acidic, low AI and	Mine water is routed into a	Often used to neutralise	Not suitable for high toxic	Nuttall and Younger (2000);
Drains (ALDs)	Fe, low dissolved	buried limestone trench which	acidity and generate alkalinity	metal mine waters; vulnerable	Watzlaf <i>et al</i> . (2000)
	oxygen concentrations	neutralises acidity and	prior to discharge to aerobic	to precipitation of AI and Fe	
		generates alkalinity	wetlands; efficient Fe and Al	on limestone; only suitable for	
			removal at low concentrations	mine waters above pH 5 with	
			(<2 mg/l)	low ferric Fe, AI (<2 mg/l) and	
				dissolved oxygen content (<1	
				mg/l)	
Oxic Limestone Drains	Net acidic, low to	An open (exposed to the	Often used to neutralise	Not suitable for high toxic	Ziemkiewicz <i>et al</i> . (1997)

(OLDs)		moderate sulphate	atmosphere)	limestone	acidity and gener	ate alkalinity	metal mine waters; high flow	
			trench which r	neutralises	prior to discharge	e to aerobic	velocities required to prevent	
			acidity and	generates	wetlands; good	rates of	Fe and AI precipitation on the	
			alkalinity		alkalinity generat	ion with low	limestone	
					water residence	times; easy		
					to construct and	d low cost		
					alternative to	o more		
					technically challe	enging and		
					costly systems			
Reducing	and	Net acidic	A layer of limestone	e beneath	Often used to	neutralise	Not suitable for high toxic	Kepler and McCleary (1994);
Alkalinity	Producing		a thick anoxic sul	bstrate of	acidity and gener	ate alkalinity	metal mine waters; requires	Jage <i>et al.</i> (2001)
Systems (F	RAPS)		organic material r	neutralises	prior to discharge	e to aerobic	significant hydraulic head	
			acidity and	generates	wetlands; efficie	nt Fe and		
			alkalinity through	processes	sulphate removal;	suitable for		
			of bacterial sulphate	reduction	net acidic mine	waters with		
			and calcite dissoluti	ion; heavy	high ferric Fe	, Al and		
			metals (mainly Fe a	nd Al) are	dissolved oxygen	content (>1		

removed as precipitates me

mg/l); low footprint

Surface Catalyzed	Net alkaline	Containers are packed with	More efficient Fe removal	Not suitable for high toxic	Younger (2000); Jarvis and
Oxidation Of Ferrous	ferruginous	high specific surface area	than aerobic wetlands; low	metal mine waters; requires	Younger (2001); Sapsford
Iron (SCOOFI)		inorganic media (e.g. plastic	footprint	significant hydraulic head;	and Williams (2009)
		trickle filter, ochre, blast		requires regular cleaning and	
		furnace slag) which		replacing of filtering media	
		encourage sorption and			
		oxidation of ferrous Fe and			
		accretion of ferric			
		oxyhydroxide			

Source control technologies and techniques

Permeable reactive Net acidic	PRBs provide a vertical and	Useful for mine waters which	Limited evidence for removal	Benner et al. (1997); Jarvis
Barriers (PRBs)	permeable compost-based of	do not emerge at the surface	of toxic metals; limited by	<i>et al.</i> (2006)
	medium in the path of a	and instead travel as	depth of aquifer	
	polluted mine water which	groundwater plumes		
	neutralises acidity and			
	promotes the generation of			
	alkalinity through bacterial			
	sulphate reduction and calcite			
	dissolution			
Physical stabilisation -	Covering of mine waste with	Immobilises contaminants at	Clay caps tend to crack in	Gandy and Younger (2003);
of mine wastes	inert material (e.g. clay, s	source and prevents	arid and semi-arid regions	Waygood and Ferriera
	gravel) to reduce oxygen	generation of mine drainage	from wetting and drying	(2009)
	inflow and water ingress into		cycles resulting in failure of	
	the contaminated material		air-tight cap	
	and, hence, the			
	concentrations of			
	contaminants in drainage			

	waters				
Chemical stabilisation -	Addition of a resinous	Immobilises contaminants at	Similar to clay caps, crusts	Tordoff <i>et al.</i> (2000)	
of mine wastes	adhesive to form a crust over	source and prevents	are prone to cracking		
	the mine waste	generation of mine drainage	resulting in failure of air-tight		
			сар		
Phytostabilisation -	A vegetative cap on the mine	Immobilises contaminants at	Concerns over bioavailability	Mendez and Maier (2008);	
	waste to immobilise	source and prevents	of contaminants to wildlife;	Pollmann <i>et al.</i> (2009)	
	contaminants by adsorption	generation of mine drainage;	need for metal tolerant plants		
	and accumulation in the	creates wildlife habitat			
	rhizosphere				
Phytoextraction -	A vegetative cap on the mine	Immobilises contaminants at	Concerns over bioavailability	Ernst (2005)	
	waste to immobilise	source and prevents	of contaminants to wildlife		
	contaminants through	generation of mine drainage;	inage;		
	hyperaccumulation in plant	creates wildlife habitat; offers			
	tissues	the possibility of recovery of			
		metals from plant tissues;			
		improves land for agriculture			

and forestry use

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