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1 Metabolic adaptation of a *Chlamydomonas acidophila* strain isolated from acid

2 mine drainage ponds with low eukaryotic diversity

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21 Abstract

The diversity and biological characteristics of eukaryotic communities within acid mine 22 drainage (AMD) sites is less well studied than for prokaryotic communities. Furthermore, for 23 24 many eukaryotic extremophiles the potential mechanisms of adaptation are unclear. This 25 study describes an evaluation of eight highly acidic (pH 1.6 - 3.1) and one moderately acidic 26 (pH 5.6) metal-rich acid mine drainage ponds at a disused copper mine. The severity of 27 AMD pollution on eukaryote biodiversity was examined, and while the most species-rich site 28 was less acidic, biodiversity did not only correlate with pH but also with the concentration of 29 dissolved and particulate metals. Acid-tolerant microalgae were present in all ponds. 30 including the species Chlamydomonas acidophila, abundance of which was high in one very 31 metal-rich and highly acidic (pH 1.6) pond, which had a particularly high PO₄-P 32 concentration. The C. acidophila strain named PM01 had a broad-range pH tolerance and 33 tolerance to high concentrations of Cd, Cu and Zn, with bioaccumulation of these metals within the cell. Comparison of metal tolerance between the isolated strain and other C. 34 35 acidophila strains previously isolated from different acidic environments found that the new strain exhibited much higher Cu tolerance, suggesting adaptation by C. acidophila PM01 to 36 37 excess Cu. An analysis of the metabolic profile of the strains in response to increasing concentrations of Cu suggests that this tolerance by PM01 is in part due to metabolic 38 39 adaptation and changes in protein content and secondary structure.

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41 Keywords: Chlamydomonas acidophila, acid tolerance, metal tolerance, acid mine
42 drainage, bioremediation, copper, zinc, cadmium

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44 **1. Introduction**

Many freshwater bodies worldwide are highly acidic either due to natural causes or
anthropogenic activities such as mining (Schultze, 2013; Smucker et al., 2014). Acid mine
drainage (AMD) due principally to pyrite oxidation, is the cause of significant acidity in lakes
and ponds situated in areas impacted by mining, and in rivers receiving mine water

49 discharge (Johnson, 2003; Nordstrom, 2000). Because decreasing pH causes increased solubility of metals, AMD results in high concentrations of dissolved Fe, S and various trace 50 metals such as Cu, Cd and Zn in the contaminated waters. Concentrations of nutrients, 51 especially inorganic phosphate (PO_4 -P), are also frequently very low (Nixdorf et al., 1998). 52 53 The combination of toxic metals and nutrient limitation limits biodiversity and can cause 54 significant ecosystem damage (Deneke, 2000; Smucker et al., 2014). Evaluation of the biological impacts of AMD allows quantification of pollution damage, allows understanding of 55 56 fundamental processes of adaptation and can identify AMD-tolerant species that have biotechnological applications, such as bioremediation (Nancucheo and Johnson, 2011; Yun 57 58 et al., 2014).

59 While prokaryotes in AMD environments has been extensively studied and reviewed 60 (Johnson and Hallberg, 2003; Mendez-Garcia et al., 2015), there is still limited knowledge 61 regarding the presence and roles of eukaryotes in these aquatic environments (Aguilera et al., 2006; Baker et al., 2004; Nixdorf et al., 1998). Photosynthetic microorganisms are found 62 in many AMD ecosystems; however, the biodiversity of phytoplankton in such waters is 63 severely limited and dominated by just a few acid-tolerant genera, such as Chlamydomonas, 64 65 Dunaliella, Euglena and Ochromonas (Aguilera et al., 2006; Hargreaves et al., 1975; Nancucheo and Johnson, 2012; Nixdorf et al., 1998; Pedrozo et al., 2001). Despite being 66 able to tolerate the highly acidic and metal-rich conditions, productivity of these extremophile 67 microalgae is often limited by low inorganic carbon and nutrient availability in acidic waters 68 (Beamud et al., 2007; Spijkerman et al., 2007b). A fairly broad diversity of heterotrophic fungi 69 and protists has also been observed in acidic waters (Baker et al., 2004; Das et al., 2009), 70 while the diversity and abundance of zooplankton is typically very low as most species are 71 72 unable to tolerate these environments (Deneke, 2000).

The high concentrations of dissolved metals in AMD can cause toxicity to
microorganisms through a wide variety of mechanisms, some of which are shared between
metals and across different organisms, such as competition with essential metals, direct
interactions with proteins and other molecules within the cell, and induction of oxidative

77 stress (Sharma and Dietz, 2009). Metals such as Cu are particularly efficient at inducing the formation of reactive oxygen species (ROS) in contrast to non-redox active metals such as 78 Zn and Cd (Valko et al., 2005). In most photosynthetic organisms, excess Cu has many 79 80 detrimental effects with the photosynthetic apparatus, including direct inhibition of 81 photosynthetic activity and degradation of chloroplast structures (Bernal et al., 2006; Küpper 82 et al., 2003). Furthermore, non-extremophile microalgae exposed to high Cu conditions exhibit high concentrations of ROS and subsequent ROS-induced damage including lipid 83 84 membrane peroxidation (Jamers et al., 2013; Jiang et al., 2016; Sabatini et al., 2009).

85 The adaptive mechanisms by which eukaryotic microorganisms including extremophile microalgae can survive in acid and metal rich conditions are still poorly 86 87 researched but potential insights into these mechanisms are increasing. For example, 88 proteomic approaches have indicated the importance of metal and acidity tolerance proteins, 89 such as molecular chaperones of the Heat Shock Protein family (Cid et al., 2010; Gerloff-Elias et al., 2006). Likewise, genome sequencing and transcriptomics studies are beginning 90 91 to identify the array of genes that might explain extremophile functional characteristics, some 92 of which may have been obtained by horizontal gene transfer from bacteria. Genome 93 sequences of the acidophiles Chlamydomonas eustigma (Hirooka et al., 2017) and Galdieria sulphuraria (Schönknecht et al., 2013) have recently been determined. Furthermore, 94 transcriptomic approaches are beginning to provide insight into the molecular mechanisms 95 of Chlamydomonas acidophila tolerance in response to Cd and Cu exposure (Olsson et al., 96 2015; Puente-Sánchez et al., 2018), and Dunaliella acidophila in response to Cd (Olsson et 97 al., 2017; Puente-Sánchez et al., 2016), although further experimental analyses of these 98 transcriptomic datasets are needed. 99

AMD tolerant biota might have potential for bioremediation, with biological-based processes potentially more cost effective and sustainable than chemical based methods such as anoxic limestone drains and chemical addition (Geller and Schultze, 2013; Hedin et al., 2010; Johnson and Hallberg, 2005). Bioremediation methods can include utilisation of bacterial SO₄ reduction and neutralisation (Neculita et al., 2007) or aerobic wetlands that can

oxidise and precipitate dissolved metals (Dean et al., 2013). However, eukaryotic algae that
can tolerate AMD conditions may be an alternative bioremediation agent (Abinandan et al.,
2018; Das et al., 2009). Novel extremophile algal strains that show high acid and metal
tolerance, and metal bioaccumulation traits are therefore needed for such applications. In
addition, extremophile algae may have other biotechnological applications, such as a source
of novel high-value chemicals including nutritional vitamins and anti-oxidants, food additives,
and biofuels (Varshney et al., 2015).

The aim of this study was to identify eukaryotes, especially extremophile microalgae, in a series of standing waters affected by AMD with the intention to characterise a strain of microalgae for evidence of AMD adaptation. Following a screen of eukaryotic biota within nine Cu-rich AMD ponds, an extremophile chlorophyte microalgal strain identified as *C*. *acidophila* was examined in detail due to its abundance and ubiquity across the site and its high tolerance to acidity and dissolved metal concentrations, especially to Cu.

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119 2. Materials and Methods

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121 2.1. Study site

The site for this study is Parys Mountain, a disused Cu mine, in Anglesey North Wales, UK. 122 The site has been mined for Cu from the Bronze Age, until mining activities ceased in the 123 early 1900s (Dean et al., 2013). The area consists of large amounts of exposed spoil, with 124 large pits and depressions that have filled with rainwater, and now retain large amounts of 125 metal-rich and acidic water (Fig. 1). In addition, precipitation ponds and lagoons were 126 constructed at the base of Parys Mountain, which were built in order to extract metals from 127 the water as part of the mining process, and also contain large volumes of AMD polluted 128 water (Younger and Potter, 2012). All the ponds are situated at close proximity within a 129 similar geology with the rocks naturally rich in Cu, Pb and Zn. It is the only known example of 130 Kuroko type volcanogenic massive sulfides in the UK, though the geology has been 131

disturbed by many millennia of underground and surface mining activities (Younger andPotter, 2012).

Of the various mining ponds and lagoons at the Parys Mountain site, nine ponds 134 were examined (Fig. 1). Ponds 1 - 4 are located at an elevated position on the spoil outcrop, 135 136 with one of these (Pond 4) on the side of a steep incline of the now drained large opencast (Fig. 1). The remaining five larger ponds (Ponds 5 - 9) are the precipitation ponds and 137 lagoons at the base of the Parys Mountain outcrop, each adjacent to agricultural land. Ponds 138 139 1 - 4 and 9 are shallow and less than 1 m depth and subject to rapid variation in depth due 140 to seasonal evaporation and rainfall. Ponds 5 - 8 are deeper and are typically 2 m in depth. All ponds showed little variation in depth across each pond. Ponds 1, 2 and 4 had no 141 vegetation in or surrounding them, whereas the other ponds had surrounding vegetation and 142 marginal wetland plants. 143

144

145 2.2. Field site sampling

Sampling at the nine AMD ponds was carried out in 2013 to 2015, including a spring 146 (February and March), summer (June) and autumn (October) sampling regime in 2015. 147 148 Water chemistry samples were taken in triplicate at each pond on each sampling occasion and were taken at approximately 15 cm depth 1 - 2 m from the edge of the pond. Water pH, 149 conductivity, temperature and dissolved oxygen were measured using a YSI 556 probe 150 (Xylem Analytics). For analysis of dissolved water chemistry metals 250 mL of pond water 151 was filtered through a 0.45 µm cellulose acetate filter, as described previously (August et al., 152 2002; Boult et al., 1994), and a 50 mL volume was retained for the analysis of dissolved 153 nutrients (PO₃-P, NO₃-N, NH₄-N). This dissolved fraction will also include some colloidal 154 metals (Florence et al., 2016). A further 50 mL volume was acidified to 1% (v/v) nitric acid 155 final concentration for the analysis of dissolved metals (AI, As, Cd, Cu, Fe, Mn, Pb, S, Zn). 156 157 The filter paper was retained, and used for the determination of particulate metals. For the analysis of algae 250 mL of unfiltered water was collected and aliquots were 158

preserved with Lugol's iodine for enumeration of algae cells and further unfiltered water

sample was taken in a sterile container for the isolation and identification of algae. Algal
samples were taken from all ponds, including samples taken from where algal biofilms were
observed (Ponds 5 and 6). For chlorophyll-*a* measurement, 250 mL of pond water was
filtered onto a GF/C filter paper.

Sediment samples were also taken to a depth of approximately 2 cm depth for determination of acid-extractable metals. For invertebrate sampling, sediment samples were taken from the littoral, approximately 1 m from the pond edge and 20 – 50 cm depth depending on the pond and sieved to remove organisms. This was followed by a 3 min sweep using a hand-held net with a mesh size of 1 mm, and a 3 min examination under large stones. Ethanol (70%) was added to preserve the invertebrate biota for identification to family level using standard keys (Greenhaigh and Ovenden, 2007; Quigley, 1977).

171

172 2.3 Nutrient, chlorophyll and metal analysis

Dissolved PO₃-P, NO₃-N, and NH₄-N were measured from the 0.45 µm filtered water 173 samples using a Skalar Sans Plus autoanalyser. Chlorophyll-a concentrations were 174 determined by absorbance spectroscopy following extraction in 96% (v/v) ethanol, as 175 176 described previously (Dean et al., 2010). Dissolved metals were determined by inductively coupled plasma atomic emission spectroscopy (ICP-AES) using a Perkin-Elmer Optima 177 5300, exactly as described previously (Dean et al., 2013). Acid extractable sediment metals 178 and suspended particulate metals were determined by acid digestion of 0.1 g of 250 µm 179 sieved dried sediments and digestion of the pond water filter papers, respectively. 180 Sediments and filter papers were digested in 5 mL ultrapure-grade nitric acid at 70°C for 4 h, 181 diluted to 2% (v/v) nitric acid, and the metals measured by ICP-AES. Certified Reference 182 Standard TM25.5 was used for all ICP-AES analyses. All samples were calibrated using a 183 matrix-matched serial dilution of Specpure multi-element plasma standard solution 4 (Alfa 184 Aesar) set by linear regression. Only results with a relative standard deviation < 20% were 185 186 considered.

187

188 2.4. Microalgae cultivation and analysis

Microalgae was visually identified to genus level and enumerated from the Lugol's iodine 189 preserved water samples by light microscopy using a Sedgewick-Rafter cell counting slide 190 and a morphological taxonomy key (John et al., 2002). Isolation of individual algae strains 191 192 was carried out by incubating serial dilutions of water samples on modified acid medium (MAM) agar plates at 22 °C with a 16-h light:8-h dark light regime and a photon flux of 193 approximately 150 µmol m⁻² s⁻¹. MAM is a defined inorganic medium developed previously 194 (Olaveson and Stokes, 1989), and used here as described by the Canadian Phycological 195 Culture Centre, University of Waterloo, Canada containing 0.5 g L⁻¹ (NH₄)₂SO₄, 0.01 g L⁻¹ 196 CaCl₂.2H₂O, 0.5 g L⁻¹ MgSO₄.7H₂O, 0.3 g L⁻¹ KH₂PO₄, 0.03 g L⁻¹ NaCl, 0.01 g L⁻¹ 197 Na₂EDTA.2H₂O, 4.98 mg L⁻¹ FeSO₄.7H₂O, 2.86 mg L⁻¹ H₃BO₃, 1.81 mg L⁻¹ MnCl₂.4H₂O, 0.22 198 mg L⁻¹ ZnSO₄.7H₂O, 0.39 mg L⁻¹ NaMoO₄.2H₂O, 79 µg L⁻¹ CuSO₄.5H₂O, 49.4 µg L⁻¹ 199 $Co(NO_3)_2.6H_2O$, 1 µg L⁻¹ vitamin B12, 1 µg L⁻¹ biotin, and 0.2 mg L⁻¹ thiamine-HCl, adjusted 200 201 to pH 3.0. Individual colonies were then extracted and grown in liquid MAM. 202 For identification of isolates using 18S rRNA gene amplification, DNA was extracted using an UltraClean Tissue & Cells DNA isolation kit (Mo Bio) and 18S rRNA gene amplicon 203 204 sequences amplified using universal primers EukF and EukR (DeLong, 1992). DNA amplicons were purified (Qiagen PCR Purification kit) before sequencing (to give ~1 kb 205 sequence reads) by GATC-Biotech, with subsequent sequence analysis performed by 206 BLAST, using the NCBI GenBank database (Table S1). The species identification of putative 207 C. acidophila strain was further confirmed by partial length 18S rDNA gene amplicon 208 sequencing, using PCR primers (18SFOR 5'-WAC CTG GTT GAT CCT GCC AGT-3', and 209 18SREV 5'-GAT CCT TCY GCA GGT TCA CCT AC-3') and PCR conditions as described 210 (Huss et al., 1999), and sequenced as described above but with sequence reads of ~1.7 kb 211 size. Phylogenetic analysis was then performed essentially as described previously 212 (Osundeko et al., 2013). 18S rRNA nucleotide sequences of selected unicellular Chlorophyta 213 214 microalgae of the Chlamydomonas moewusii clade were obtained from GenBank and 215 sequences were aligned using ClustalW. The phylogenetic tree was generated using the

maximum likelihood method using RAxML-GUI and the GTR-GAMMA model (Stamatakis,
2006). Confidence in the tree was assessed using the thorough bootstrap method by
performing 10 runs of 100 replications.

219 An isolate of C. acidophila from Pond 1 (named PM01) was cultivated to quantify its 220 tolerance to high metal concentrations and pH ranges alongside two strains of C. acidophila 221 previously isolated from an acidic pond near Fratiskovy Lazne, Czech Republic (CCAP 11/136) (Fott and McCarthy, 1964), and an acidic mining Lake 111 in eastern Germany 222 223 (CCAP 11/137) (Gerloff-Elias et al., 2005). All cultures were maintained in MAM pH 3.0. 224 Cultures were serially inoculated in fresh media and were grown in batch culture conditions on an orbital shaker at 120 rpm at 22 °C with a 16-h light:8-h dark light regime and a photon 225 flux of approximately 150 µmol m⁻² s⁻¹. For metal exposure treatments, various 226 concentrations of metals as chloride salts were added to liquid MAM pH 3.0 or solid MAM pH 227 228 3.0 agar plates. For pH range treatments, MAM was adjusted and buffered to the desired pH (from pH 1.0 to 7.0) as described previously (Gerloff-Elias et al., 2006). Essentially the 229 230 medium was buffered through the presence of Fe in the medium for pH 1.0 to 3.0, with 10 mM citric acid for pH 4.0 to 5.0, and with 10 mM HEPES for pH 6.0 to 7.0. The pH could be 231 232 maintained to within ~0.5 pH unit during the growth period. Starting cell densities were normalised by optical density measurement at 680 nm (OD_{680nm}). Algae growth was 233 determined by cell number, and in some instances by OD_{680nm}, total chlorophyll or growth 234 rate measurement, exactly as described previously (Osundeko et al., 2013). Algae samples 235 were prepared for metal content measurement by ICP-AES, as described previously 236 (Webster et al., 2011). Cells were EDTA washed to remove externally bound metals, as 237 validated elsewhere for Cd, Cu, Pb and Zn (Hassler et al., 2004). Cells were centrifuged for 238 10 min at 3000 g, followed by resuspension of the cell pellet in 10 mL of 1 mM EDTA for 5 239 min, then re-centrifuged and washed with a 15 mL volume of Milli-Q water. After further 240 centrifugation, cell pellets were oven-dried at 60 °C for 24 h and then digested in 0.5 mL of 241 ultrapure concentrated nitric acid at 70 °C for 3 h. Samples were diluted in Milli-Q water to 242 243 2% (v/v) concentration of acid and analysed by ICP-AES as described above. Cells of C.

244 *acidophila* were measured (width and length) using an eyepiece graticule, and cell volume 245 was calculated according the prolate spheroid formula (Hillebrand et al., 1999). The cell 246 volume measurements were used to calculate the internal (cellular) concentration of metal 247 (determined from EDTA-washed cells) on a volume basis. This value was then divided by 248 the external (MAM or pond water) concentration of metal in order to calculate concentration 249 factor (K_{conc}) values.

250

251 2.5. Fourier transform-infrared (FT-IR) spectroscopy

252 Each algal strain was grown in liquid MAM (pH 3.0), with Cu concentrations of 0, 6.5, 13 and 130 mg L^{-1} for 14 days, at which point cultures had an OD_{750nm} of between 0.3 and 0.5. 253 254 Cultures were normalised to an OD_{750nm} of 0.3 and 10 mL was centrifuged at 1800 g for 5 min. The supernatant was removed and the cells washed in 1.5 mL of 0.9% (v/v) NaCl, then 255 256 centrifuged and washed again before final resuspension of the cell pellet in 1 mL of 0.9% (v/v) NaCl before 20 µL of this suspension was deposited onto a 96-well silicon microplate. 257 The samples were then oven dried at 40°C for 1 h, and an additional 20 µL sample was 258 added and the samples once again dried at 40°C for 1 h. The plate was placed in a HTS-XT 259 260 high-throughput microplate extension and FT-IR spectra collected using an Equinox 55 FT-IR spectrometer (Bruker Corporation), equipped with a deuterated triglycerine sulphate 261 detector. Spectra were collected over the wavenumber range 4000-600 cm⁻¹. Each sample 262 was analysed as nine technical replicates. Spectra were pre-processed using extended 263 multiplicative signal correction (Martens and Stark, 1991) prior to multivariate analysis. Band 264 assignments were determined as described previously (Driver et al., 2015). 265

266

267 2.6. Modelling and statistical analysis

268 Metal speciation modelling was performed using Visual Minteq version 3.0 (Gustafsson,

269 2010). All data was statistically analyzed by one-way ANOVA using Tukey post-hoc test

- 270 performed using Prism v.6.04 (GraphPad). Principal component analysis (PCA) of
- 271 environmental data was performed using PRIMER v.6 (Primer-E) and plotted using XLSTAT

(Addinsoft), while PCA of FT-IR spectra was performed using MATLAB version R2016a. All
 environmental data (except for pH values) were natural log transformed for PCA and linear
 regression analysis.

275

276 **3. Results**

277

278 3.1. Chemical characteristics of extremely acidic metal-rich mining ponds

279 The nine ponds were situated in close proximity (Fig. 1) but differed in water chemistry (Fig. 280 2). In all ponds, concentrations of suspended particulate metals were present at much lower 281 concentrations compared to dissolved metals (Fig. S1). None of the metal or nutrient 282 concentrations in the ponds differed significantly across the spring, summer and autumn 283 samples. Ponds 1, 2 and 4 were situated on top of the Parys Mountain site in depressions 284 of the spoil waste. These ponds were characterised by very low pH (< 2.0) and high conductivity values (Fig. 2A). Pond 1 in particular was highly acidic (mean pH of 1.6) and 285 286 had a very high conductivity (mean 10.03 mS), which is predominantly controlled by the high dissolved S concentration, as well as high concentrations of dissolved Fe, Al, Cd, Cu and Zn 287 288 (Fig. 2B, C). In addition, Pond 1 also had a very high PO₄-P concentration (Fig. 2D) and the highest dissolved As concentration (Fig. S1). Ponds 2 and 4 also had high dissolved metal 289 and S concentrations but differed from Pond 1 due to lower conductivity (~3 mS) and much 290 lower PO₄-P. Pond 3 was also situated on the top of the mine site, however, this pond stood 291 out as the least polluted, with a weakly acidic pH (mean pH 5.3), and low conductivity and 292 dissolved metals and S (Fig. 2A; Fig. S1). The precipitation ponds and lagoons (Ponds 5 -293 9) were also highly acidic but had lower conductivity due to much lower dissolved metal 294 concentrations, with the exception of dissolved Mn (Fig. S1). In most ponds, ammonium 295 (NH₄-N) was the dominant form of N rather than nitrate (NO₃-N), as is expected in acidic 296 297 waters where nitrification rates are low (Baffico et al., 2004). Pond 6 had unusually high NO₃-N (Fig. 2D) possibly because the pond is directly adjacent to fertilised farmland. 298 299 Chlorophyll-a was detectable in all ponds, indicating the presence of photosynthetic

microorganisms, but Pond 1 also differed from the other ponds with respect to having
significantly higher concentration of chlorophyll-*a*, which increased substantially during the
year as the water temperature and insolation increased (Fig. 2E).

The differences in water chemistry between the ponds situated within the mine spoil 303 304 waste (Ponds 1, 2 and 4), the lagoons and precipitation ponds (Ponds 5 - 9), and the less polluted Pond 3, are reflected in the PCA plot (Fig. 3). In particular, Pond 1 and Pond 3 were 305 distinguished from each other and from the other ponds through difference in pH. 306 307 Furthermore, the high sediment AI and Mn concentrations explained the differentiation of 308 Pond 3, likely a consequence of the high pH of this water resulting in the precipitation of 309 dissolved AI. Pond 1 was situated within the PC space particularly on the basis of high 310 chlorophyll-a, PO₄-P and dissolved As concentration. It is therefore apparent that many of these ponds are highly toxic environments, particularly Pond 1 with its very high acidity and 311 312 high concentration of toxic dissolved trace metals including As, Al, Cd, and Cu.

313

314 3.2. Taxa diversity

Overall taxa diversity, as determined by number of different taxonomic families, was lowest in the highly acidic, high conductivity Pond 1 (with two taxa) and also low in the Ponds 2 and 4, which are also situated among open mine spoil. The highest number of taxa were found in the low conductivity, mildly acidic (pH 5.3) Pond 3, with 15 taxa, while the

lagoons/precipitation ponds had intermediate taxa diversity (7-10 families). Linear regression 319 analyses of the relationships between taxa diversity and water chemistry (Fig. S2) showed a 320 significant positive correlation between increasing pH values and increasing taxa number (R²) 321 = 0.59; p = 0.03) but also a very strong negative correlation between decreasing conductivity 322 and increasing taxa number ($R^2 = 0.85$; p = 0.001), indicative of the dissolved metal 323 concentration. In particular, there was significant correlation between dissolved As, Cd, Cu, 324 325 Fe and S and taxa number, as well as between particulate Fe and S with taxa number, but 326 no significant correlation on the basis of sediment metal concentration (Fig. S2).

327 Invertebrates were observed in all ponds, though only Chironomidae was identified in all, although there were only one or two individuals recorded from Pond 2 and 4 (Fig. 5A). In 328 Ponds 1 and 4, this was the only invertebrate taxa present, indicating that the conditions in 329 330 these ponds are not suited for a diverse invertebrate community, but are able to support a 331 few species that have adapted to the extreme AMD conditions. Other invertebrate taxa that 332 were abundant are Corixidae, present in five of the ponds, and Sialidae, present in four of 333 the ponds, but both absent in the three most acidic ponds (Fig. 4). The observed biota in 334 Pond 3 included 14 invertebrate families, including many that are pollution-sensitive. Ponds 335 5 – 9 contained between 5 and 7 invertebrate taxa. All ponds were devoid of macrophytes within the open water. 336

337 A number of distinct eukaryotic microorganisms were isolated from the open water including microalgal, fungal and protozoan species (Table S1). The highest eukaryotic 338 diversity was in Ponds 2, 5 and 6 (Fig. 4). In Ponds 5 and 6 and their adjacent channels 339 much of the microorganisms were observed to be associated in biofilms. These included a 340 strain which showed 99% identity based on the 18S rRNA sequence to Euglena mutabilis 341 (Table S1), a well-known acid-tolerant species that has been previously observed in AMD 342 343 environments, including coal mine waste sites (Brake et al., 2001), and including at the adit draining from the Parys Mountain mine (Nancucheo and Johnson, 2012). In addition, a non-344 motile chlorophyte (likely to be Koliella corcontica; 96% identity) that has been previously 345 observed in mildly acidic lakes (Vrba et al., 2003), and a diatom (likely to be Eunotia 346 naegelii; 98% identity) was found associated with biofilm adjacent to Pond 6. However, 347 diatoms were not found in open water in any of the Parys Mountain ponds. Ponds 1, 3 and 7 348 had the lowest number of eukaryotic microorganisms, with just one chlorophyte algae taxa 349 350 identified in each pond (Fig. 4). There was substantial heterogeneity between the ponds, 351 with all microbial taxa present in just one pond sample with the exception of the Euglena sp. present in two ponds, and a *Chlamydomonas* sp. identified in all ponds (100% abundance). 352 353 Sequencing of the ~1 kb 18S rRNA gene amplicon from the Chlamydomonas sp. showed 354 highest sequence identity (99%) to a strain annotated as C. acidophila (CCAP 11/134)

(Table S1). Quantification of cell density of this strain found highest density in Pond 1 (Fig. 5A), which correlated significantly with the highest pond PO₄-P concentration ($R^2 = 0.60$; p = 0.02), but cell density was also lowest in the most alkaline pond water where there was the lowest dissolved metal concentrations, likely explaining the positive correlation between cell density and conductivity or dissolved metal concentrations, such as Cu (Fig. 5B).

360

361 3.3. Phylogeny of a *C. acidophila* strain

362 C. acidophila was found across the Parys Mountain site in all nine ponds and was highly 363 abundant in the most acidic and metal rich Pond 1. Therefore this strain was studied further in more detail. The isolate of *C. acidophila* from Pond 1 (named PM01) was examined by 364 365 longer (1.7 kb) 18S rRNA gene read sequence and phylogenetic analysis (Fig. 6). PM01 18S 366 rRNA sequence was identical apart from one nucleotide within this 1.7 kb region to 367 sequences from three strains annotated as C. acidophila (Gerloff-Elias et al., 2005): CCAP 11/134 (an isolate from Argentina), CCAP 11/136 (an isolate from Czech Republic) and 368 CCAP 11/137 (an isolate from Germany). PM01 was also identical within this 18S region to 369 an unidentified strain (Rt1n1) originally isolated from the acidic Rio Tinto river in Spain 370 371 (Amaral Zettler et al., 2002). The CCAP 11/136 strain (Fott and McCarthy, 1964), previously regarded as an authentic strain of C. acidophila (Gerloff-Elias et al., 2005), is also deposited 372 in a different culture collection as strain UTCC 354 (Pollio et al., 2005), but this has distinct 373 18S rDNA sequence (Fig. 6), indicating that CCAP 11/136 and UTCC 354 are not in fact 374 identical. It was previously argued that the UTCC 354 strain assigned as C. acidophila may 375 be more appropriately assigned as Chlamydomonas pitschmannii, another highly 376 acidotolerant species (Pollio et al., 2005). Other strains recorded as C. acidophila including 377 UTCC 121 (an isolate from Canada) (Twiss, 1990) and OU 030/a (an isolate from Japan) 378 (Nishikawa and Tominaga, 2001), were also distinct from PM01 and CCAP 11/136 but 379 380 grouped more closely to the UTCC 354 strain and C. pitschmannii (Fig. 6). Although strain 381 OU 030/a has also been previously regarded as an authentic strain of *C. acidophila* 382 (Nishikawa and Tominaga, 2001; Pollio et al., 2005), here we refer to the CCAP 11/136 and

11/137 strains as *C. acidophila* species, in line with previous analysis (Gerloff-Elias et al.,

2005), and thus the PM01 strain is further referred to as a strain of this species.

385

386 3.4. AMD tolerance by *C. acidophila* PM01

387 To examine the relationship between pH and C. acidophila further, strain PM01 from Pond 1 was grown in MAM artificial pond water at a range of pH values from pH 1.0 to 7.0. PM01 388 exhibited a very broad pH tolerance range, with optimal growth at pH 3.0 - 5.0 (mean growth 389 rate ranging between 0.158 – 0.175 d⁻¹; no significant difference between treatments; p >390 0.05), and with the highest cell density after 25 d obtained in pH 3.0 conditions. Strong 391 growth was still observed in pH 7.0 (mean growth rate 0.153 d⁻¹) and pH 2.0 (mean growth 392 rate 0.157 d⁻¹) conditions, but with a significant reduction (p < 0.05) in growth after 25 d in pH 393 394 2.0 and pH 7.0 by 25% and 37%, respectively compared to pH 3.0 MAM. and could grow at pH 1.0 after an extended lag phase of 13 d (growth rate 0.018 d⁻¹), but with a significant 395 reduction in growth by 73% after 25 days compared to pH 3.0 MAM. The pH characteristics 396 397 of the ponds may explain in part the microalgae biodiversity and cell density profiles, and as described above, there was a significant negative correlation ($R^2 = 0.57$; p = 0.02) between 398 399 pH and C. acidophila cell density (Fig. 5B).

Samples of PM01 taken directly from Pond 1 showed high concentration of absorbed 400 401 and internalised metals, as determined by measurement of EDTA-washed cells, to remove externally cell wall-bound metals. There was a high concentration of Cu and Zn accumulated 402 almost entirely within the cell (no significant difference between EDTA washed versus 403 unwashed cells) (Table 1). Relative to mean Pond 1 dissolved Cu concentration of 58.6 mg 404 L⁻¹, the internal cellular Cu concentration in pond 1 cells was 84.3 fg cell⁻¹ (equivalent to 405 185.5 mg L⁻¹). The mean concentration of dissolved Zn in Pond 1 was 37.9 mg L⁻¹, while the 406 internal cellular Zn concentration was 115.0 fg cell⁻¹ (253.0 mg L⁻¹). This gives concentration 407 factor (K_{conc}) values of 3.2 for Cu and 6.7 for Zn. Substantial bioconcentration was also 408 observed for Pb and Cd. Relative to a mean Pb concentration in Pond 1 of 0.8 mg L⁻¹, the 409 410 accumulation and bioconcentration of Pb by the strain was particularly high, with a cellular

411 concentration of 146.3 fg cell⁻¹ (321.8 mg L⁻¹) giving a K_{conc} value of 421.8. Cd was also 412 accumulated almost entirely within the cell to a concentration of 2.8 fg cell⁻¹ (6.2 mg L⁻¹), and 413 relative to pond water concentration of 0.3 mg L⁻¹ gives a K_{conc} value of 24.5. In contrast, 414 only 56% of accumulated Fe was taken up into the cell (Fig. S3).

415 Metal tolerance and accumulation by C. acidophila PM01 was further examined in an artificial growth medium to assess the tolerance range of three of the trace metals found 416 within the Parys Mountain ponds. PM01 was grown in increasing concentrations of Cd, Cu 417 418 and Zn in MAM at pH 3.0. For cells grown under controlled conditions, the maximum cell 419 density achieved and the total chlorophyll concentration per cell (as a measure of cell physiological status) was higher than that observed for the cells analysed in situ (in Pond 1) 420 (Table 1). The PM01 strain displayed tolerance to high concentrations of these three metals. 421 To allow comparison with Pond 1 water concentrations, the free ionic Cd²⁺, Cu²⁺ and Zn²⁺ 422 concentrations were calculated using the Visual Minteg speciation model. PM01 could 423 tolerate up to 2.6 mg L⁻¹ Cd²⁺, which was 16-times higher than present in Pond 1 water, with 424 425 no significant inhibition of growth rate, maximum cell density or chlorophyll content when compared to no added Cd^{2+} (Table 1). However, at a concentration of 6.0 mg L⁻¹ Cd²⁺ all 426 427 three parameters (growth rate, cell density, chlorophyll content) were significantly reduced (p < 0.05). PM01 was highly tolerant to Cu, and none of the concentrations up to 78.3 mg L^{-1} 428 Cu²⁺ treatment significantly inhibited growth or chlorophyll-a concentration, the apparent 429 reduction in cell density was not significant. Substantial Zn tolerance was also observed, 430 with no significant inhibition to growth rate at 855.4 mg L⁻¹ Zn²⁺, which was nearly 30-times 431 higher than the Zn^{2+} concentration in Pond 1, but at 1760.8 mg L⁻¹ Zn^{2+} there was a 432 significant (p < 0.05) reduction in maximum cell density, although growth rate and chlorophyll 433 concentration was not significantly inhibited. The strain was still growing in 3002.8 mg L⁻¹ 434 Zn^{2+} despite cell density being inhibited by 94%, with also a significant (p < 0.05) reduction in 435 growth rate and chlorophyll concentration (Table 1). 436

Accumulation of Cd, Cu and Zn was also quantified in the metal-treated PM01 strain
after 25 d growth in MAM pH 3.0 (Table 1). There was a concentration-dependent increase

439 in cellular accumulation of Cu and Cd with no significant difference between the values with or without EDTA washing, indicating that almost all of the Cu and Cd was taken up within the 440 cell. In contrast, there were significant (p < 0.05) differences in Zn concentration following 441 442 EDTA washing, indicating that a smaller proportion of Zn was internalised (Table 1). As the 443 Zn concentration in the medium increased, the relative concentration within the cell 444 decreased, suggesting that Zn transport into the cell saturated at higher concentrations. 445 Overall, the characteristics of metal accumulation in artificial media were broadly similar to 446 those observed in the pond.

447

448 3.5. Metabolic adaptation to Cu tolerance by C. acidophila PM01

449 It was unknown whether the metal tolerance properties of C. acidophila differ between strains isolated from AMD sites with differing water chemistry. To begin to assess this, the 450 451 tolerance of PM01 to a range of metals (AI, Cd, Cu, Fe, Mn and Zn) was compared to two other strains of C. acidophila (CCAP 11/136 and CCAP 11/137) that had previously been 452 isolated from different field sites. Strains CCAP 11/136 and CCAP 11/137 were validated by 453 18S rDNA sequencing and phylogenetic analysis as C. acidophila species (Fig. 6). The 454 455 CCAP 11/136 strain was originally isolated from a highly acidic (~pH 1.0) humic acid-rich peat water environment but the metal characteristics of the site were not described (Fott and 456 McCarthy, 1964). In contrast, strain CCAP 11/137 originated from German mining Lake 111 457 at pH 2.6, very low total P (8 μ g L⁻¹) and fairly high levels of Zn (0.75 mg L⁻¹) (Spijkerman et 458 al., 2007a). 459

For Al, Cd and Mn there was no growth difference between the three strains. However, Fe treatment substantially inhibited growth of CCAP 11/136, while Zn treatment slightly inhibited growth of PM01 and CCAP 11/137 (Fig. S4). However, there was a very clear-cut difference in growth between the three strains following Cu exposure (Fig. 7). On solid media containing 130 mg L⁻¹ Cu, the CCAP 11/137 strain was unable to grow, and the CCAP 11/136 strain grew very weakly in contrast to strong growth by the PM01 strain (Fig. 7A). In liquid media with addition of 0.0 mg L⁻¹, 6.5 mg L⁻¹ and 3 mg L⁻¹ growth rate and cell

density after 14 days was identical between all three strains. However, with 130 mg L^{-1} Cu addition, cell growth was unchanged for PM01 but both CCAP 11/136 and 11/137 strains were barely able to grow (Fig. 7B).

To examine whether there were any macromolecular changes within the strains in 470 471 response to the increasing copper treatment, and to examine whether the different strains could be distinguished on the basis of their metabolic 'fingerprint', the FT-IR spectroscopy 472 technique was used. FT-IR spectra were collect for replicates of each strain cultivated in the 473 absence of added Cu or with the addition of 6.5 and 13 mg L⁻¹ Cu (Fig. S5). In addition, 474 PM01 was tested at the 130 mg L⁻¹ concentration that inhibited growth of 11/136 and 11/137, 475 476 thereby preventing FT-IR spectroscopy analysis of these strains at the higher Cu 477 concentration. PCA of all FT-IR spectra showed that the CCAP 11/137 strain samples 478 cluster separately from CCAP 11/136 and PM01 in all treatments, with the close clustering of 479 spectra indicating that Cu addition did not significantly alter the metabolic fingerprint of CCAP 11/137 (Fig. 8A). Likewise Cu addition did not significantly alter the metabolic 480 481 fingerprint of the CCAP 11/136 strain, with all samples clustering with the PM01 control samples. However, there was a clear difference in the FT-IR spectra-derived metabolic 482 483 profile of PM01 samples following addition of increasing Cu concentrations, with the sample position within the PCA plot changing on the basis of both PC1 and PC2. At the highest 484 (130 mg L⁻¹) Cu concentration all replicate PM01 samples are distinct from the control 485 samples (Fig. 8A). The PC loading plots (Fig. 8B) indicate that along PC1 the spectral 486 changes are based predominantly on increased abundance of amide peaks at 1655 and 487 1545 cm⁻¹, as well as a decrease in one of the carbohydrate peaks at approximately 1036 488 cm⁻¹. 489

490

491 4. Discussion

This study has demonstrated that while AMD substantially impacts biodiversity in aquatic
environments, there is still substantial taxa abundance in these extreme locations. The
scarcity of invertebrates across the ponds clearly indicates the severity of pollution at this

495 abandoned mine site. With the exception of the near-neutral pH Pond 3, invertebrate diversity was very poor in all ponds. Taxa diversity overall, including invertebrate diversity in 496 497 particular, was strongly correlated with pH and with dissolved metal concentration, which 498 was in line with expectations and previous studies (Courtney and Clements, 2002; Malmqvist 499 and Hoffsten, 1999). Nevertheless, some invertebrate species can adapt to extreme AMD 500 conditions (De Bisthoven et al., 2005; Deneke, 2000), and can be used as indicators of mine-waste pollution (Gray and Delaney, 2008). Here Chironomidae were observed in all 501 502 ponds, including the very acidic Pond 1 and 4, and highly abundant in Pond 9, probably 503 because this pond is fairly shallow and has fine sediment in contrast to the other ponds. Certain Chironomidae species are able to survive in highly acidic waters, such as the acid-504 505 tolerant Chironomus acidophilus that has been found nearby in the highly acidic (pH 2.4) and 506 metal-rich Afon Goch river draining Parys Mountain (Michailova et al., 2009).

507 While Chironomidae were present in all ponds so there was also the consistent presence of chlorophyte microalgae, and in particular a strain confirmed as C. acidophila. 508 The widespread occurrence of C. acidophila at Parys Mountain reflects the extremely acidic 509 and metal-rich pond water and is consistent with other acidic and metal-rich sites (Fott and 510 511 McCarthy, 1964; Gerloff-Elias et al., 2005; Hargreaves et al., 1975; Twiss, 1990). The widely differing C. acidophila abundance between ponds may reflect nutrient availability. AMD 512 environments typically have very low productivity, due in part to low nutrients such as PO₄-P 513 (Spijkerman et al., 2007a; Spijkerman et al., 2007b), and in this study nutrients, and in 514 particular PO₄-P, were at low concentration in all but Pond 1. The low cell density of this 515 species in Ponds 2 – 9, where PO₄-P concentrations were low suggests growth limitation 516 due to low PO₄-P availability. In contrast, Pond 1 had very high PO₄-P and a high 517 concentration of *C. acidophila* cells. Other studies have shown high PO₄-P levels associated 518 with acidic lakes in areas with abundant PO₄-P-containing FeS minerals (Spijkerman, 2008); 519 however, as Pond 2 is within 5 m of Pond 1 and has the same mineralogy but significantly 520 lower PO₄-P levels, the high concentration of PO₄-P in Pond 1 is unlikely to be due to 521 522 minerology of the surrounding area. It may be that the high PO₄-P concentration is due to a

eutrophication event, as there is evidence of construction waste deposition that is likely to explain the PO_4 -P entry into the pond.

The high microalgae productivity of Pond 1 is due almost exclusively to C. acidophila 525 526 abundance. Acidophilic microalgae present in the other ponds, such as K. corcontica, E. 527 naegelii and E. mutabilis, were absent from Pond 1. E. naegelii and other acid-tolerant 528 diatoms are increasingly used as indicator species for acid polluted environments (Zalack et 529 al., 2010) and have been previously found in surface sediment from one of the rivers 530 draining from the Parys Mountain site (Dean et al., 2013), as well as in acid pit lakes with 531 equivalent water chemistry (Geller, 2013). However, their apparent low abundance in these ponds might partly be due to the sampling regime used here from the near-surface open 532 water rather than from the bottom of the ponds. E. mutabilis was observed in two of the nine 533 ponds (Ponds 5 and 6), which are both ~pH 3.0, a pH range that has been shown to be 534 535 preferential for this organism (Brake et al., 2001). Furthermore, other acidophilic microalgae such as Ochromonas sp. and Dunaliella sp., that are widely abundant in many acidic sites 536 (Aguilera et al., 2006; Nixdorf et al., 1998), were not identified in Pond 1, or in any other 537 Parys Mountain ponds. That C. acidophila was the only microalgal species identified in Pond 538 539 1 is likely to be due to the extreme conditions that restricts microalgal diversity. One of these factors appears to be water pH, and the *C. acidophila* strain studied here was particularly 540 acid tolerant. In waters with pH > 3.0, it has been previously observed that algal biodiversity 541 is generally higher (Smucker et al., 2014), yet the least acidic pond studied here, Pond 3 (pH 542 5.3), still had very low algal biodiversity and very low C. acidophila abundance, suggesting 543 that other factors are also important, such as nutrient availability or water chemistry. 544 Although the dissolved Fe, Zn, Cu and Al concentrations in Pond 1 did not exceed 545

the very high concentrations seen in some AMD lakes and rivers such as those of the lberian Pyrite Belt (Sánchez España et al., 2008), they did exceed those seen in mine pit lakes in Germany, Poland, Australia and USA (Geller, 2013). The metal concentrations of the shallow Pond 1 can thus be considered high. The strain of *C. acidophila* isolated from Pond 1 is therefore not just extremely acid-tolerant, but can also tolerate high metal

concentrations. The PM01 strain showed substantial tolerance to Cd, Cu, and Zn. A
comparison of PM01 to other genera of acid tolerant algae shows the Zn tolerance of PM01
was higher than that reported for *Chlorella protothecoides var. acidicola* isolated from AMD
sites in a Spanish mine and *E. mutabilis* isolated from the adit flowing from Parys Mountain.
However, both of these strains showed a higher Cu tolerance than PM01 (Ňancucheo and
Johnson, 2012).

Other studies looking at metal tolerance in C. acidophila are fairly scarce, though a 557 558 study looking at a putative strain of C. acidophila OU 030/a isolated from a volcanic acid lake 559 also showed high tolerance to Cd, Cu and Zn in metal-rich minimal media (at pH 4.0) when compared to other algal species (Nishikawa and Tominaga, 2001). Furthermore, a strain of 560 C. acidophila RT46 isolated from Río Tinto in Spain, showed unaffected photosynthetic 561 activity in response to 0.5 mM Cu (~32 mg L⁻¹ Cu) exposure (Olsson et al., 2015). A putative 562 C. acidophila strain (UTCC 121) isolated from Cu contaminated soil was previously shown to 563 tolerate up to 100 mg L⁻¹ Cu. However, in contrast, a laboratory strain of *C. acidophila* 564 (CCAP 11/96) was Cu sensitive, as was the non-acidophilic freshwater alga 565 Chlamydomonas reinhardtii (Twiss, 1990), indicating that the Chlamydomonas genera is not 566 567 intrinsically Cu tolerant. In this study we demonstrate that the C. acidophila strain isolated in this study (PM01) has higher Cu tolerance than strains (CCAP 11/136 and 11/137) from 568 AMD field sites with less Cu pollution. This shows that the PM01 strain has adapted to the 569 high dissolved Cu concentrations of Pond 1, rather than this species having innate Cu 570 tolerance properties. 571

572 FT-IR spectroscopy analysis demonstrates that the tolerance of PM01 to copper is 573 partly due to its ability to modulate its metabolism in response to increasing Cu exposure, as 574 indicated by a dose-dependent change in spectra characteristics, which does not occur with 575 the other *C. acidophila* strains. Examination of the FT-IR spectra indicates that this metabolic 576 adaptation is predominantly due to protein increase and potential modification of protein 577 secondary structure, as shown by significant increase in amide I peak height associated with 578 C=O stretching, and in amide II peak height associated with N-H bending and C-N stretching

579 (Giordano et al., 2001). A previous study using FT-IR spectroscopy to examine sensitivity and subsequent acclimation of microalgae to wastewater treatment, also found that the 580 acclimation process was coincident with a relative increase in amide I and amide II peak 581 582 height (Osundeko et al., 2014). Moreover, this same study showed that particularly sensitive 583 strains including Chlamydomonas debaryana and Desmodesmus intermedius exhibited 584 accumulation of carbon storage products including glycerolipids and starch, while acclimated strains that could tolerate the wastewater conditions did not show this response. Likewise, 585 586 other metabolic indicators of stress such as increased carbohydrate and lipid peaks within 587 the FT-IR spectra were not observed in response to high Cu treatment in PM01.

588 Because Cu exerts toxicity in part through inhibition of key cellular processes 589 including photosynthesis, either directly due to Cu binding or indirectly via accumulation of 590 ROS (Jamers et al., 2013; Küpper et al., 2003; Sabatini et al., 2009), it would be expected 591 that adaptive mechanisms would counteract these processes in some ways. Proteomic responses to Cu stress linked to Cu tolerance have previously been observed in other 592 organisms. Cu exposure experiments in plants and fungi have observed increases in soluble 593 protein that has been linked to induction of anti-oxidant enzymes (Cavalcanti Luna et al., 594 595 2015; Gao et al., 2008; Rout et al., 2013) while induction of Cu-binding proteins has been demonstrated in a Cu-tolerant variety of rice (Chen et al., 2015). Mechanisms of Cu 596 tolerance in algae are not well understood, and may involve differential Cu uptake and 597 internalisation in some cases (Levy et al., 2008). Although the molecular mechanisms of 598 stress tolerance by C. acidophila are also poorly understood but there have been some 599 recent insights. Heat shock proteins, which are a family of evolutionarily conserved stress 600 tolerance molecular chaperones, have been previously found to increase in abundance in C. 601 acidophila CCAP 11/137 in response to very low pH and metal-rich lake water treatment, 602 partly in response to high Fe concentration (Gerloff-Elias et al., 2006; Spijkerman et al., 603 604 2007a). To date, no proteomic or enzymatic analysis has been performed in C. acidophila in 605 response to Cu stress, but a transcriptomics approach observed differences in mRNA 606 transcript profiles following Cu treatment in C. acidophila RT46. A range of gene transcripts

were up-regulated in response to Cu treatment including those involved in photosynthesis,
signaling and stress-response (Olsson et al., 2015). Future experiments will aim to examine
the proteomic adaptive response of *C. acidophila* PM01 in more detail in order to enhance
our fundamental understanding of metal tolerance in microalgae, also to appreciate the
effects that environmental pollution has on adaptive evolution and the ecological
consequences of such adaptation.

613 A potential application of highly metal tolerant microalgae such as C. acidophila 614 PM01 is the potential use of such organisms for metal bioremediation. Indeed PM01 was 615 confirmed to bioconcentrate metals through a combination of cell wall binding and 616 internalisation. Algae strains, such as those isolated in this study, may be used for in situ 617 lake bioremediation in surface water mesocosms by controlled eutrophication and harvesting (Dessouki et al., 2005), or for ex situ bioremediation, such as immobilised algae in 618 619 bioreactors (Mehta and Gaur, 2005). The accumulated metals may then be harvested and processed to allow metal recovery (Minoda et al., 2015; Raikova et al., 2016). Alternatively, 620 highly acid and metal tolerant microalgae such as the PM01 strain, may have an important 621 role in sustaining SO₄-reducing bacteria by providing organic carbon and thus increasing the 622 623 efficiency of AMD remediation microbial bioreactors (Diez-Ercilla et al., 2014; Nancucheo and Johnson, 2012; Totsche et al., 2006). For example, it was demonstrated that microalgal 624 addition to mine tailing mesocosms containing pyrite-oxidizing bacteria caused higher 625 production of alkalinity, higher concentrations of ferrous Fe, and increased immobilization of 626 Cu and Zn (Nancucheo and Johnson, 2011). 627

The main aim of this study was to identify microorganisms from the open, standing waters of the AMD ponds. Some of the ponds were surrounded by vegetation including wetland plant species, which will harbor associated microorganisms (Aguinaga et al., 2018). Although not the scope of this study, future research can examine the role of the plants on microbial communities within isolated AMD pond environments, as well an examination of biota along spatial transects of the ponds including within the sediment. Moreover, future studies will be needed to examine how other microorganisms found in these environments

have adapted to AMD stresses, and whether there are common mechanisms betweendifferent extremophile species.

637

638 **5. Conclusions**

639 AMD is a major source of freshwater pollution worldwide for which restoration is very 640 important. However, as found in this study, while extremely acidic and metal rich AMD 641 substantially impacts biota, there is still substantial biodiversity, with tolerance derived 642 through natural adaptation. In particular, a strain of *C. acidophila* is abundant in all ponds at 643 this Cu mine site, especially in waters with high acidity and coupled with high PO_{4} -P 644 concentration. Although Cu toxicity is a significant challenge to most photosynthetic 645 organisms, this strain of C. acidophila has specifically adapted to the high Cu status of the ponds in contrast to other strains of the same species isolated from field sites elsewhere. 646 647 Moreover, the C. acidophila strain displays evidence of Cu-dependent metabolic plasticity. The marked metal tolerance and metal accumulation characteristics of C. acidophila PM01 648 indicates that organisms from these environments have biotechnological potential, such as 649 bioremediation. 650

651

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921 Figures



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- 923 Fig. 1. Map and satellite image of Parys Mountain copper mine in Anglesey, North Wales,
- 924 UK. The sampled AMD ponds are labelled 1–9 and the AMD-polluted Southern Afon Goch
- 925 river is indicated. The location coordinates of each sampled pond are also shown (A). (B)
- Satellite image of the mine and the sampled ponds. Source, Google Earth, 2016.



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929 Fig. 2. Water chemistry of the AMD ponds. (A - C) pH and conductivity values (A), and concentration of dissolved Fe, Cd (B), Cu and Zn (C) for each pond during 2015 (spring, 930 summer, autumn samples). Boxes show the 25th and 75th percentiles, the line within the 931 boxes shows the median values, and the whisker bars show minimum and maximum values 932 (n = 3 - 18). (D) PO₄-P, NO₃-N and NH₄-N concentration in each pond (June sample). Pond 933 934 7 data is not available. Values are means (n = 3 - 12) and error bars correspond to the 935 standard error of the mean. (E) Mean chlorophyll-a (Chl a) concentration in each pond during 2015 (spring, summer, autumn samples). Chl a change in Pond 1 during the year and 936 pond water temperature is shown (inset). Each pond was sampled in triplicate on three 937 separate occasions (spring, summer, autumn). Values are means (n = 9) and error bars 938 939 correspond to the standard error of the mean. For all data, bars that do not share a lower 940 case letter show significant difference (p < 0.05) between pond sites. 941



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Fig. 3. PCA of water chemistry data from each of the AMD ponds. (A) Observation plot for each pond. The upper site small ponds are shown as blue triangles, the near-neutral pH pond as a red circle, and the lower site large lagoon ponds as green squares. (B) Variables plot with selected environmental variables labelled.



Fig. 4. Taxa diversity of eukaryotic organisms in the AMD ponds as determined by numbers of distinct taxonomic families of microalgae, moss/liverwort, fungi, protozoa, as identified by 18S rRNA gene amplicon sequencing, and families of invertebrates, as identified by visual

952 observation and use of taxonomic identification keys. Data for Pond 8 is not available.



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Fig. 5. (A) Abundance of *C. acidophila* microalgae and Chironomidae insects in each pond 955 956 from summer sampling. Values of *C. acidophila* are means (n = 3) and error bars correspond 957 to the standard error of the mean, while value of Chironomidae are total counts. For C. acidophila data, bars that do not share a lower case letter show significant difference (p < 958 959 0.05) between pond sites. Chironomidae data for Pond 8 is not available. (B) Linear regression analyses for dissolved phosphate, pH, conductivity and dissolved Cu in relation to 960 C. acidophila cell counts in AMD ponds. Apart from pH values, all data were natural log 961 962 transformed.





Fig. 6. 18S rDNA sequence analysis of C. acidophila strains. Phylogenetic tree based on 965 966 18S nucleotide sequence obtained from the PM01 strain isolated from Pond 1, sequences of C. acidophila CCAP 11/136 and CCAP 11/137 strains, other known or putative C. acidophila 967 strains, and selected unicellular Chlorophyta microalgae of the Chlamydomonas moewusii 968 clade, including three strains originally classified as C. acidophila. The sequence lengths 969 were between 1528 - 1792 nucleotides. Chlamydomonas reinhardtii is included as an out-970 group. Accession numbers are shown for each sequence. Bootstrap percentage values are 971 indicated at the tree nodes of branches for 100 replications and indicate confidence in tree 972 node positions. The branch length scale bar indicates evolutionary distance. 973



Fig. 7. Cu tolerance of *C. acidophila* PM01 in comparison to *C. acidophila* strains CCAP 11/136 and 11/137. (A) Growth of algae spot dilutions on MAM pH 3.0 plates with or without added Cu and photographed after 12 d. Image is representative of 3 independent experiments. (B) Growth of strains in liquid MAM pH 3.0 determined after 14 d cultivation in response to a range of Cu concentrations. Values are means (n = 3 - 4) and error bars correspond to the standard error of the mean. Bars that do not share a lower case letter show significant difference (p < 0.05) between strains.

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Fig. 8. PCA clustering of FT-IR spectra from *C. acidophila* PM01 in comparison to *C. acidophila* strains CCAP 11/136 and 11/137 in response to a range of Cu concentrations. (A) PCA scores plot of replicate (n = 4) FT-IR spectra obtained from cells grown after 14 d cultivation in liquid MAM pH 3.0 with increasing concentrations of added Cu or no added Cu. Only PM01 cells could grow in 130 mg L⁻¹ Cu. (B) PC1 and PC2 loading plot. Band peaks which explain most of the variation for each PC are ν C=O of amide I (1655 cm⁻¹) and δ N-H of amide II (1545 cm⁻¹).