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Distinct Microbial Communities in the Soils, Waters and Speleothems of a Hyperalkaline Cave System

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Key Points:

- Samples from soils, waters, calcites and cave muds were examined to understand the distribution of microbiota in a hyperalkaline cavern.
- The microbial biomass in above-cave soils and cave muds was notably higher than that in speleothem calcites and drip waters.
- Limited in-cave microbial similarities between microhabitats indicated a high variability likely driven by the steep alkalinity gradient.

22 **Abstract**

23 Caves are complex ecosystems with various microbial habitats. Understanding the
24 individual community structures in the key source environments (soils, waters) and
25 those in the sinks (speleothems, cave muds) can help elucidate the extent to which in-
26 cave communities are a function of their source communities, or if they can
27 successfully adapt and diversify locally. Here, we assess the network of microbial
28 communities existing within a unique British cave system. Poole's Cavern is
29 characterised by alkalinity or even hyperalkalinity (pH>9) in its drip waters, creating
30 a series of challenging ecological niches for microbes to survive. Samples of soil, drip
31 water, speleothem and cave sediment were collected for chemical characterization,
32 cell culture assays and DNA sequencing analysis. We show that microbial abundance
33 and diversity were highest in the soils proximal to the cave, and the microbiotas of the
34 hyperalkaline cave chamber were less abundant and less diverse than the cave
35 chamber with lower pH. Proteobacteria and Planctomycetota were the most prevalent
36 bacterial phyla throughout the cave system irrespective of pH, indicating their
37 members are comparatively more metabolically versatile. Predicted essential
38 metabolic pathways were still dominant within all sampling sites, and microorganisms
39 were shown to be capable of utilising various inorganic or simple organic compounds
40 to survive. Interestingly, co-occurrence between Poole's samples was limited, never
41 demonstrating more than 50% similarity. This work highlights the diversity of
42 microbial communities within this extreme environment and the development of
43 microbial niches which reflects the adaptation strategy of microorganisms under
44 alkaline-hyperalkaline conditions.

45 **Plain Language Summary**

46 Caves are complex environments, forming a range of living spaces for microbes.
47 Dissolved inorganic minerals and organic matter are carried by water that flows from
48 the soils above caves to speleothems and into sediments on the ground. Poole's
49 Cavern in the UK is a cave system with very high pH (>9) waters. These waters are
50 not friendly to most microbes living in neutral pH environments and thus possibly
51 foster special microbial communities that can stay alive under high pH conditions.
52 Full of calcite formations and without observable fauna, this cave is not rich in
53 nutrients. We sampled soils above Poole's Cavern, drip waters, speleothems and cave
54 muds to determine the biomass and microbial species in them. We found that the soils
55 outside of this cave had the highest microbial content and diversity. The cave chamber
56 with higher pH waters had fewer microbes than that with lower pH waters. We
57 demonstrate that connectivity of microbial species between in-cave sampling sites
58 was limited, and the community of microbes in each site developed relatively
59 independent finding ways to survive in their respective environment. This study
60 presents the high potential of microbes to evolve and adapt even in conditions almost
61 inhospitable to life.

62 **1 Introduction**

63 Cave and karst systems represent some of the least studied extreme
64 environments on the planet. Caves are in most cases characterised by extreme nutrient
65 deficiency and utter darkness (Dong et al., 2020), characteristics quite inhospitable to
66 life. However, understanding the karst critical zone (Chen et al., 2018) is of
67 fundamental importance to vast numbers of the global population who rely on karst
68 aquifers for fresh drinking water supplies (Stevanovic, 2019). Possibly the least well-

69 established connections within karst and cave science revolve around the role of
70 microbial communities within karst, their sources, abundance, community structure
71 and their ability to adapt and even thrive in cave environments (Engel, 2010). It is
72 possible that microbial communities play a central role in the karst aquifer system;
73 their utilisation and consumption of key nutrients (C, N, S and P) may help to balance
74 the anthropogenic increase in nutrient loads to many karst environments. With
75 microbes holding such a potentially critical role within the karst geochemical system
76 and wider ecosystem, it is fundamental that we tackle key questions surrounding the
77 “microbial network” (Zhu et al., 2021).

78 Previous work on cave microbiology is scarce but slowly developing
79 (Fairchild and Baker, 2012; Zhu et al., 2021). The routes to microbial deposition
80 within cave systems are thought to primarily be through the entrainment of microbes
81 along with other organic matter in the soil and the “wash in” of this material through
82 the karst aquifer and into the cave. There is also the potential for aquifer biofilms to
83 develop and contribute to communities later deposited within the cave system. In
84 some systems, cave dwelling animals or human visitors could play a role in adding to
85 the microbial network possibly through aerosol transport and contamination (Dredge
86 et al., 2013; Smith et al., 2013), although this is thought to diminish in importance as
87 you move further from the cave entrance and away from tourist routes (Fairchild and
88 Baker, 2012).

89 Based on DNA studies, it has been noted that about half of all recognised
90 bacterial phyla have been identified within cave or karst ecosystems (Engel, 2010;
91 White, 2009) and that microbial communities can thrive in these ecosystems with cell
92 counts of up to 10^6 cells/g of rock (Barton and Jurado, 2007). Globally, studies have
93 identified Betaproteobacteria, Deltaproteobacteria, Acidobacteria, Actinobacteria, and
94 Nitrospira throughout cave and karst aquifers, associated in-cave sediments and on
95 cave surfaces, including speleothems (Farnleitner et al., 2005; Ortiz et al., 2013;
96 Pronk et al., 2009; Shabarova and Pernthaler, 2009). The lack of sunlight within cave
97 systems means that these microbes have to be adapted to utilise alternative energy
98 sources. For example, the chemosynthetic fixation of inorganic carbon, degradation of
99 aromatic hydrocarbons and the use of hydrogen sulphide have all been identified as
100 non-photosynthetic energy pathways in use in cave systems (Marques et al., 2019;
101 Sarbu et al., 1996; Wu et al., 2015). These microbial communities may become the
102 primary food base to support other cave adapted species (Sarbu et al., 1996). Not only
103 are these species highly adapted to the cave environment, they also appear specifically
104 adapted to their niche within the cave. A detailed study of caves of the Yunnan-
105 Guizhou Plateau (China) found that cross-cave microbial networks were often similar,
106 however individual “in-cave” habitat niches (air, water, rock and sediment) were
107 found to be quite distinct from one another (Zhu et al., 2019; Zhu et al., 2021),
108 potentially controlled by their different source micronutrients (Cloutier et al., 2017)
109 and mineral chemistry (Wu et al., 2015).

110 In an attempt to undertake a comprehensive assessment of the microbial
111 network and its key niches within an anthropogenically impacted cave system, we
112 consider possible microbial habitats from the top down, looking at community
113 dynamics within soils, karst aquifer water, sediments and calcite speleothem deposits.
114 This work has focussed upon Poole’s Cavern, Buxton, UK. This site was chosen
115 primarily due to the unique hyperalkaline drip waters (pH>9) found within some
116 sections of the cave, the small amount of previous microbial work undertaken locally,
117 and the availability of baseline cave climate data provided as part of the British Cave

118 Science Centre (BCSC) project. This cave system offers an excellent opportunity to
119 interrogate microbial dynamics in a high pH environment, investigate the unique
120 development of microbial niches and quantify the extent to which there are linkages
121 between these habitats.

122 Whilst most cave and karst systems have very little known about their inter-
123 niche microbial network, the unique hyperalkaline soil and karst water environments
124 around Buxton have attracted slightly more previous work on extremophile
125 communities. Burke et. al. (2012) profiled a distinct anaerobic alkaliphilic community
126 dominated by a single, unidentified bacterial species within the *Comamonadaceae*
127 family of Betaproteobacteria in the Buxton lime kiln deposits near the Poole's Cavern
128 site. This community appears capable of microbial nitrate reduction with increasing
129 anoxia (Burke et al., 2012). There is also evidence of communities surviving in a non-
130 karst hyperalkaline spring in Buxton. Smith et. al. (2016) sequenced the community in
131 pH 7.3 to pH 13 waters. At pH 13, the sequence library was dominated by the families
132 *Pseudomonadaceae* and *Enterobacteriaceae* of Gammaproteobacteria, with low
133 overall diversity. They concluded that these communities were functioning at a pH of
134 11-12, but at higher pH (pH 13) these communities remained present, but were
135 unlikely to be active. The only previous work within Poole's Cavern focussed on soil
136 and hyperalkaline drip waters. Blyth et. al. (2014) conducted a study looking at lipid
137 biomarkers: glycerol dialkyl glycerol tetraethers (GDGTs) (Blyth et al., 2014). They
138 found two distinct profiles between the subsurface soils above the cave and the
139 speleothem drip waters in an interior chamber. The drip waters were dominated by
140 branched GDGTs, indicative of a bacterial source, and the soils were dominated by
141 isoprenoid GDGTs, indicative of an aquatic archaeal source. This difference indicates
142 that microbial communities established within the cave (or karst) were distinct from
143 those in the soil zone, suggestive of microbial diversification within the karst system,
144 rather than a simple in-wash from infiltrating soil waters. These previous local studies
145 offer a glimpse into the potential communities that can exist within hyperalkaline
146 waters such as those sampled in Poole's Cavern.

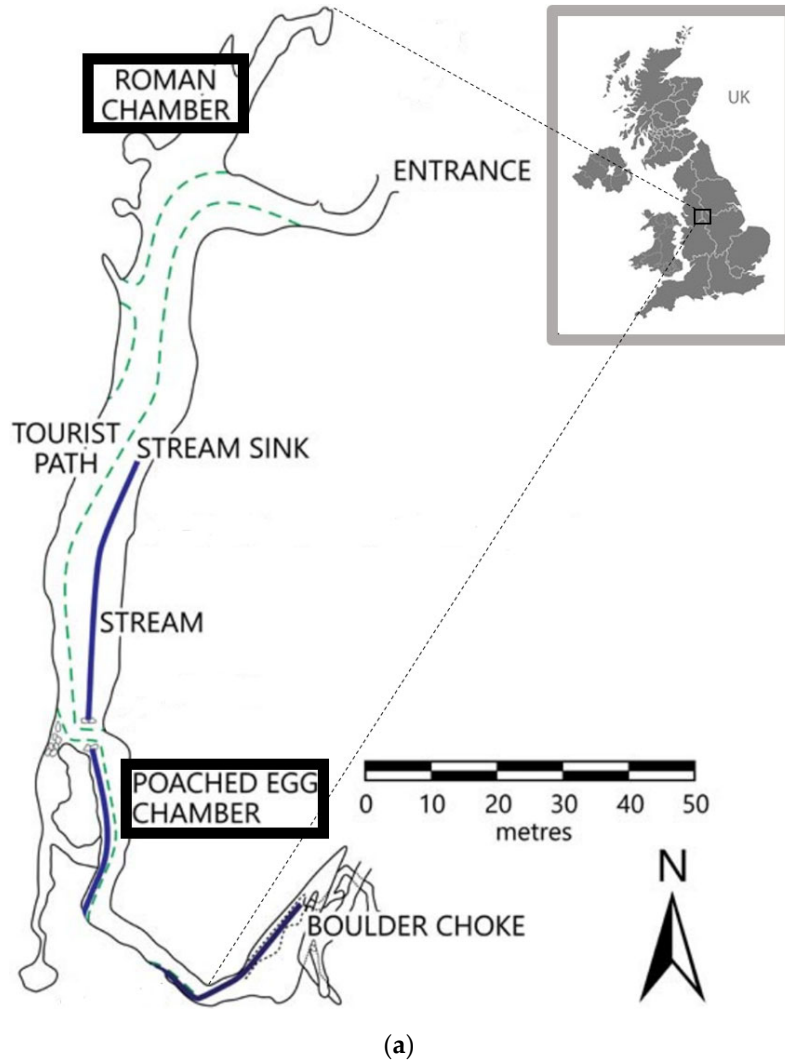
147 The interior environmental conditions of Poole's Cavern vary remarkably.
148 However, the cavern is a cross-linked integrative ecosystem. Unveiling the individual
149 communities in the key source environments (soils, waters) and those in the sinks
150 (speleothems, muds) will elucidate the development of above-cave and in-cave
151 biodiversity and the adaptive capacity of microorganisms to different but proximal
152 microhabitats (Zhu et al., 2021). This paper will investigate the community network
153 more fully, using culture-dependent experiments and DNA sequencing to answer
154 questions about community viability and structure throughout the soil, karst aquifer,
155 cave and speleothem system.

156 **2 Site Description and Methodology**

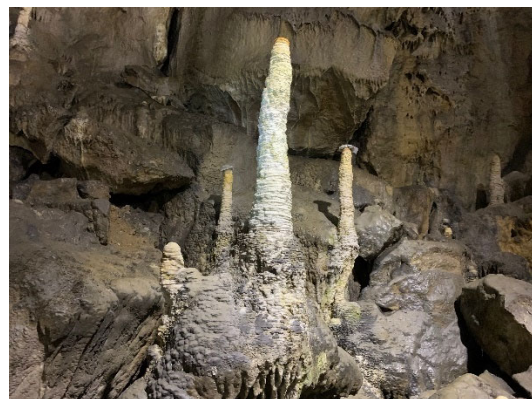
157 2.1 Site description

158 Poole's Cavern is a shallow epigenetic cave evolved from early Carboniferous
159 Bee Low limestones (Hartland et al., 2010b; Newton et al., 2015). The main cave strikes
160 N-S with a total length of 240m, with the main passage terminated by a partially
161 cemented boulder choke (Deakin et al., 1968; Rowberry et al., 2020). The main
162 chambers of interest for sampling (Figure 1a) from this project include Roman Chamber
163 (RC, close to the cavern entrance) and Poached Egg Chamber (PE, deeper into the
164 cavern) (Figure 1b, c). These two chambers are both out of reach from the tourist path

165 and the public (Figure 1a). Speleothem formations in the PE chamber are characterised
166 by extremely rapid growth rates (10 mm/year) (Hartland et al., 2010b), caused by the
167 super-saturation of carbonate under hyperalkaline conditions. This is due to the
168 leaching of 17th Century lime kiln waste (CaO) above the cave producing Ca(OH)₂,
169 which rapidly disassociates to Ca²⁺ and OH⁻ ions raising the pH (Hartland et al., 2010b).
170 At hyperalkalinity, the hydroxylation of CO₂ dominates, producing a rich carbonate
171 source (Clark et al., 1992) that has facilitated the formation of the large distinctive
172 stalagmites that attract tourists annually (Figure 1c).



(b)



(c)

173 **Figure 1. (a)** Locations of Poole’s Cavern and its chambers of this study (modified
174 with permissions, from Rowberry et al., 2020). Internal landscapes of **(b)** Roman
175 Chamber (RC) and **(c)** Poached Egg Chamber (PE) [BGS © UKRI].

176 Above Poole’s Cavern exists a well-established, managed deciduous
177 woodland. Soils are organic rich with significant deciduous leaf litter overtopping a
178 ~10-cm layer of organic-rich topsoil and a ~30-cm layer of lime kiln waste (Charles et
179 al., 2015; Hartland et al., 2010a). This lime kiln waste is left over from several lime
180 kilns, which were atop some of the cave chambers. As waters leach the kiln waste
181 they become hyperalkaline. Where leaching waters enter the cave system they form
182 distinctive stalagmite deposits in the chambers below.

183 No vertebrates including bats were observed in the Poole’s Cavern due to the
184 adverse in-cave conditions as well as the mesh on the entrance door. The closure of
185 the cave to tours (more than one year before sampling) during the COVID-19
186 pandemic provided a unique opportunity to sample the cave in a more natural state,
187 where any microbiological impact from anthropogenic interference is limited.

188 2.2 Poole’s Cavern and sample collection

189 In total, eight samples were targeted, including soil, water, sediment and
190 calcite. Sampling took place in spring/summer (April) 2021. Two calcite samples (PE
191 High and PE Low) were “grown” in Poached Egg Chamber. These samples were
192 allowed to precipitate onto clean, autoclaved watch glasses, with calcite left collecting
193 for two months before sampling (Figure 1a). One sample (PE Straw) was from a
194 stalactite straw on the ceiling of the PE Chamber. A PE drip water sample (PE Water)
195 was collected over a period of two weeks by attaching a funnel to an actively
196 depositing speleothem (‘PE3’, as described in (Morgan, 2022)) and routing the
197 captured water into a 20-L jerrican. 6 L of this water sample were filtered through a
198 0.22 µm Sterivex filter within Poole’s Cavern and immediately sealed with Parafilm.
199 RC chamber water was collected only for a single water chemical measurement due to
200 slow drip rates. Additionally, one calcite and one mud sample (RC Calcite and RC
201 Mud) were acquired from Roman Chamber (Figure 1b), with the mud being extracted
202 from the bottom of a seasonally active pool fed by drip waters. This pool contained
203 water at the time of sampling but is known to dry out during the summer months.
204 Moreover, soils above Poole’s Cavern and calcareous soils from ~1.7-km southeast of
205 the nearby Brook Bottom valley (53°14’15.2”N, 1°54’35.8”W) were sampled. All
206 samples for microbiological analyses (sequence analysis and cell count analysis) were
207 collected using a sterilised hand auger and containers. For soil and sediment samples,
208 the top 10-cm layers were discarded before sample collection; each soil or sediment
209 sample was a pool of at least three locations.

210 All samples for microbiological analysis were transported on ice to BGS
211 Keyworth that day. Samples for DNA analysis were frozen at -20°C on return.
212 Samples for cell culture were shipped cold to the University of St Andrews and stored
213 at 4°C before processing.

214 Temperature, pH, and electrical conductivity of speleothem drip waters from
215 PE and RC chambers were measured *in situ* at the time of sampling using a WTW
216 Multi 340i handheld multimeter.

217 2.3 DNA extraction and sequence analysis

218 Samples for sequence analysis were frozen after sampling and stored at -20°C

219 for one week before DNA extraction. All implements for molecular analyses were
220 either filter sterilised, autoclaved, flamed, or UV-irradiated to prevent any external
221 contamination. The de-ionised water used was molecular biology grade and nuclease-
222 free. Either ~400 mg of solid material or the Sterivex filter membrane were used as
223 input for DNA extraction. 100 µL DNA were extracted with one negative control using
224 the FastDNA SPIN Kit for Soil (MP Biomedicals, CA, USA) following the
225 manufacturer's instructions. DNA extracts were amplified for barcoded MinION 16S
226 metagenomic sequencing using 16S Barcoding Kit (SQK-RAB204; Oxford Nanopore
227 Technologies, Oxford, UK) and analysed using Flongle R9.4.1 flow cells (Oxford
228 Nanopore Technologies, Oxford, UK). The 16S rRNA primer pair used for
229 amplification was the domain Bacterial-specific forward primer 27F (5'-
230 AGAGTTTGATCCTGGCTCAG-3') and the universal reverse primer 1492R (5'-
231 GGTTACCTTGTTACGACTT-3') (Reysenbach et al., 2000). The polymerase chain
232 reaction (PCR) was conducted using C1000 Touch thermocycler (BioRad, CA, USA)
233 with the following conditions – 95°C for 1 min, then 40 cycles of 95°C for 20 s, 55°C
234 for 30 s and 65°C for 2 min, with final extension at 65°C for 5 min. PCR reactions were
235 conducted in triplicate, each consisted of 5 µL LongAmp Taq DNA polymerase (NEB,
236 MA, USA), 0.2 µL barcoded primer, and 0.03-1.6 µL DNA extract, with the remaining
237 volume made up with molecular grade water. The barcoded PCR products were purified
238 using ChargeSwitch PCR Clean-Up Kit (Thermo Fisher Scientific, MA, USA).
239 Amplicon concentration was quantified using an Invitrogen Qubit 3.0 Fluorometer.
240 Only samples with more than 0.05 ng/µL yields (all samples other than PE High and
241 Low) were passed for sequencing on the Oxford Nanopore MinION platform (Oxford
242 Nanopore Technologies, Oxford, UK).

243 Bacterial sequences were analysed in the framework of the open-source
244 program Quantitative Insights into Microbial Ecology 2 (QIIME 2 version 2021.8)
245 (Bolyen et al., 2019). In the first step, the MetONTIIME meta-barcoding pipeline
246 (version 1.14.0) was applied as an EPI2ME 16S workflow emulator to process
247 MinION sequenced fast5 data in QIIME 2 framework (Maestri, 2021; Maestri et al.,
248 2019). Next, base calling was performed using Guppy 5.0.11, followed by read
249 dereplicating and clustering at 100% identity (Maestri, 2021) against the Silva 138
250 99% reference database (Quast et al., 2013; Yilmaz et al., 2014). Then, the generated
251 amplicon sequence variants (ASVs) were aligned against the Silva 138 99% reference
252 database to output taxonomy data using q2-feature-classifier (Bokulich et al., 2018). To
253 pledge the sequence quality, sequences with confidence > 0.7 were selected for
254 taxonomic identification (Table S1). Based on the taxonomy information,
255 mitochondrial, chloroplast, archaeal, eukaryotic, and unassigned ASVs were excluded
256 prior to the taxonomic classification analysis. Then, the alpha diversity index, Faith's
257 phylogenetic diversity, was computed after rarefaction at sampling depth 92 with
258 alpha-phylogenetic package (Faith, 1992) in QIIME 2.

259 The improvement of QIIME 2's genome prediction will subsequently optimize
260 accuracy of metabolic inference. Thereafter, potential functions of these 16S
261 sequences were predicted with Phylogenetic Investigation of Communities by
262 Reconstruction of Unobserved States 2 (PICRUST2) (Douglas et al., 2020) based on
263 individual ASVs. PICRUST2 has also remarkably improved accuracy and flexibility for
264 marker gene metagenome inference compared to its predecessor. ASVs at this stage
265 were 3,982 counts. Briefly, reads were assigned into phylogenetic units with HMMER
266 (Eddy and Wheeler, 2007), EPA-ng (Barbera et al., 2019), and GAPPA (Czech and
267 Stamatakis, 2019). The hidden-state prediction was performed with the R package

268 ‘caster’. Representative ASVs with nearest-sequenced taxon index (NSTI) more than
269 two were excluded from the output. Finally, microbial metabolic pathways were
270 identified using the Minimal set of Pathways (MinPath) approach (Ye and Doak,
271 2009) referencing the KEGG Orthology (KO) and MetaCyc pathway database, with
272 the correlation coefficients around 0.8-0.9 between the predicted and observed data
273 (Douglas et al., 2020). Please note the PICRUST2 inference method is difficult in
274 predicting reduced genomes (e.g., rare species) and cannot perfectly reflect active or
275 even present genes from extracted microbiome, but provides a most likely function
276 based on DNA sequences and related well-understood species (Douglas et al., 2020).
277 After PICRUST2 filtering, 22 (8 from PE Water, 4 from Soils above Poole’s, 9 from
278 Brook Bottom soils, 1 from RC Calcite) out of 3977 sequences were removed from
279 functional group analysis. These 22 sequences belonged to 5 genus *Candidatus*
280 *Omnitrophus* (Verrucomicrobiota), 1 genus *Brevundimonas* (Alphaproteobacteria), 1
281 genus *Hydrogenophaga* (Gammaproteobacteria), 1 unidentified Proteobacteria, and
282 14 unknown bacteria (Table S1).

283 2.4 Determination of microbial abundance

284 The estimated cell content of each sample was calculated from the
285 concentration of extracted DNA using the equivalent conversion factor 9×10^{-15} g
286 DNA·cell⁻¹ (Kirchman, 2012).

287 Additionally, microscopic cell counting and cell culture of aseptically
288 collected samples (stored at 4°C) were performed one week after sampling. Each
289 sample was suspended and well-mixed in 1:1 volume (mL):weight (g) autoclaved
290 0.9% NaCl solution with two drops of Tween 20 to preserve osmotic pressure and
291 detach cells from particles. Several 10× serial dilutions were further prepared. Cell
292 numbers of 100:1 dilutes were counted under an AmScope optical microscope using a
293 Hirschmann hemocytometer in quadruplicate; each replicate picked up the large
294 counting grids at four corners.

295 For cell culturing experiments, the required volume of each dilute sample was
296 estimated and spread on Plate Count Agar culture plates based on preliminary
297 culturing tests. The recipe for Plate Count Agar was 5 g/L tryptone, 2.5 g/L yeast
298 extract, 1 g/L dextrose and 9 g/L agar. Each sample was plated on these culture plates
299 in quadruplicate. Visible colonies were counted after 10 days of incubation at 21°C in
300 accordance with the growth rate of these microbiotas.

301 2.5 Statistical techniques

302 Hierarchical clustering of species from each individual site were performed
303 using the Bray-Curtis dissimilarity index and the unweighted pair group method with
304 arithmetic mean (UPGMA) algorithm in Past 4.08 (Hammer et al., 2001). Next,
305 bivariate correlation coefficients were calculated to understand the relationship
306 between sites. Prior to correlation analysis, normality test was performed to determine
307 whether parametric (Pearson’s r) or nonparametric (Kendall’s tau) statistical approach
308 should be used. Since the microbial community network closely links with ecosystem
309 function (Finlay et al., 1997), we conducted network analysis on Poole’s Cavern
310 microbiota at species level. Samples were combined for the extended Bayesian
311 information criterium graphic lasso (EBICglasso) estimation (Friedman et al., 2008).
312 Using individual sites and intercorrelation as nodes and connecting edges
313 respectively, the co-occurrence pattern of species that represents the microbial
314 network was defined based on their EBICglasso coefficients. The network diagram

315 and centrality plot of expected influence were drawn using JASP 0.16.2 ([https://jasp-](https://jasp-stats.org/)
316 [stats.org/](https://jasp-stats.org/)).

317 **3 Results**

318 3.1 Water chemistry

319 The temperatures of PE Water and RC Water were 7.3 ± 0.3 °C (n=4) and 9.3
320 °C (n=1), respectively. Electrical conductivities of each water type were 1216.5 ± 57.5
321 $\mu\text{S}/\text{cm}$ (n=4) and $600.0 \mu\text{S}/\text{cm}$ (n=1), respectively. pH values were 12.2 ± 0.1 (n=5)
322 and 9.3 (n=1), for PE Water and RC Water respectively.

323 3.2 Microbial abundance and diversity

324 Sites where samples were collected for microscopic cell counting and cell
325 culture experiments were: PE High, PE Low, PE Water, PE Straw, Soils above
326 Poole's, and Brook Bottom soils. PE Low had the lowest detectable microbial content
327 as reflected in $<0.05 \text{ ng}/\mu\text{L}$ and in all cell count methods; the extracted DNA
328 concentration from PE High was also below the detection limit of Qubit fluorometer
329 ($0.05 \text{ ng}/\mu\text{L}$), although its countable microbes were more abundant than PE Water and
330 PE Straw. Conversely, the microbial abundance of Soils above Poole's Cavern and at
331 Brook Bottom, were consistently the highest (Table 2).

332 **Table 1**

333 *Environmental Parameters of PE and RC Chambers within Poole's Cavern.*

Features	PE	RC
pH	12.2 ± 0.1 (n=5)	9.3 (n=1)
Temperature (°C)	7.3 ± 0.3 (n=4)	9.3 (n=1)
Conductivity ($\mu\text{S}/\text{cm}$)	1216.5 ± 57.5 (n=4)	600.0 (n=1)
Hydroxide (mg/L)	144.4 (n=1)	-
Carbonate (mg/L)	11.6 (n=1)	-
Total alkalinity (mg/L)	156.0 (n=1)	-
Orthophosphate (mg/L)	0.018 ± 0.001 n=3)	-

334 *Note.* RC chamber did not have enough liquid sample for environmental
335 characterization.

336 **Table 2**

337 *The Concentrations (Mean \pm Standard Error) of Microbial Cells Determined by Plate*
338 *Count Agar Cell Culture, Microscopic Cell Counting, and Soil DNA Conversion;*
339 *Faith's Phylogenetic Diversity Indices of Poole's Cavern Samples.*

Samples	Viable Cultivable Cell Counts (CFUs/g)	Microscopic Cell Counts (cells/g)	Soil DNA Equivalence (cells/g)	Faith's Phylogenetic Diversity
PE High	$(2.36 \pm 1.67) \times 10^4$	$(1.28 \pm 0.50) \times 10^8$	$< 1.39 \times 10^6$	-
PE Low	$(6.00 \pm 1.47) \times 10^2$	$(3.44 \pm 1.64) \times 10^7$	$< 1.39 \times 10^6$	-
PE Water	$(5.13 \pm 2.81) \times 10^3$	$(2.81 \pm 1.43) \times 10^7$	5.89×10^7	16.21
PE Straw	$(2.41 \pm 1.53) \times 10^4$	$(2.50 \pm 1.29) \times 10^7$	2.40×10^7	17.68
Soils above Poole's	$(1.29 \pm 0.24) \times 10^7$	$(4.14 \pm 1.11) \times 10^8$	1.61×10^9	24.35
Brook Bottom soils	$(7.83 \pm 2.72) \times 10^5$	$(1.83 \pm 0.78) \times 10^8$	1.61×10^9	26.22
RC Calcite	-	-	4.36×10^7	22.49
RC Mud	-	-	2.73×10^8	22.12

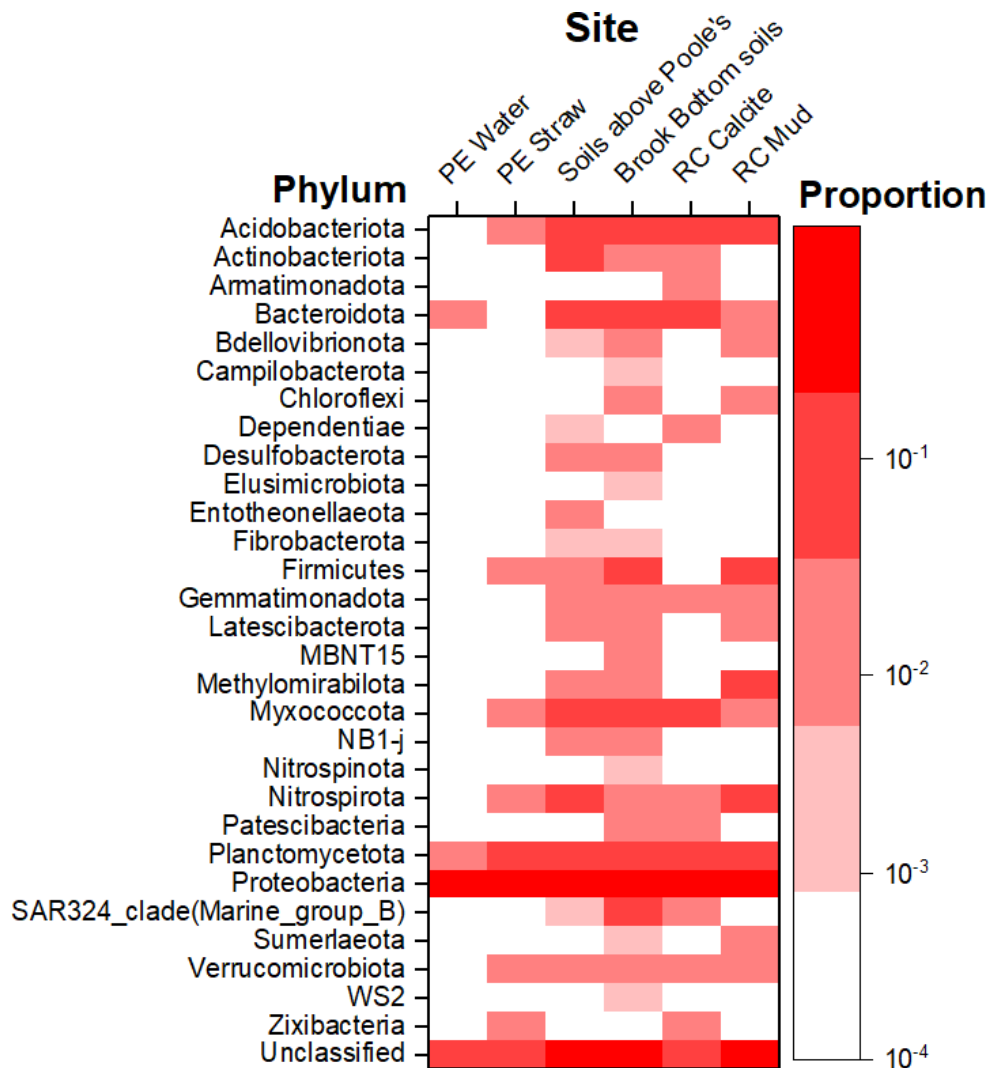
340 *Note.* RC Calcite and Mud did not have enough sample for cell culture. PE High and
 341 Low did not have enough extractable DNA for sequencing.

342 Based on sequence analysis, PE Water had the lowest phylogenetic diversity
 343 index (16.21), followed by PE Straw (17.68). Brook Bottom soils (26.22) and Soils
 344 above Poole's (24.35) had the highest diversity indices. The two RC samples, calcite
 345 (22.49) and mud (22.12) were intermediate to these (Table 2).

346 3.3 Taxonomic phylogeny

347 Proteobacteria were the most abundant phyla and displayed similar relative
 348 abundance among six samples (Figure 2). Proportions of unclassified bacteria were
 349 high in all samples. PE Water had abundant Actinobacteriota and Bacteroidota, while
 350 PE Straw had higher abundances of Firmicutes and Zixibacteria (Figure 2). Consistent
 351 with phylogenetic diversity indices, PE Water and PE Straw had the least diverse
 352 communities while soils from Brook Bottom and above Poole's had the most diverse
 353 communities (Table 2). Planctomycetota, Verrucomicrobiota, Acidobacteriota,
 354 Myxococcota and Nitrospirota were abundant in the solid samples (mud, soil, calcite),
 355 while PE Water clearly had lower abundances of these groups (Figure 2).

356



357

358 **Figure 2.** Proportions of taxonomic groups of bacteria from Poole's Cavern samples.

359 The dominant bacterial families displayed very different patterns among
360 Poole's Cavern sampling sites (Table 3). The most dominant family was always from
361 Proteobacteria however differed among the samples: *Xanthomonadaceae* was the
362 most dominant family in PE Water (28.6%), which also included high abundances of
363 *Comamonadaceae* (26.1%). *Comamonadaceae* was the most dominant in PE Straw
364 (16.3%), Soils above Poole's (5.8%), and RC Calcite (8.3%); *Nitrosomonadaceae*
365 was the most dominant in Brook Bottom soils (10.8%) and RC Mud (14.1%). Other
366 dominant families came from Planctomycetota, Nitrospirota, Bacteroidota,
367 Methyloirabilota, Firmicutes, Sumerlaeota, Acidobacteriota and Zixibacteria (Table
368 3).

369 **Table 3**370 *Proportions of Dominant Bacterial Families (>2% in At Least One Site, Bolded) from Poole's Cavern Samples.*

Family	PE Water	PE Straw	Soils above Poole's	Brook Bottom soils	RC Calcite	RC Mud
<i>Nitrosomonadaceae</i> (Proteobacteria)	0.8%	11.4%	2.9%	10.8%	6.7%	14.1%
<i>Sphingomonadaceae</i> (Proteobacteria)	12.6%	0.0%	1.1%	0.0%	3.3%	0.0%
<i>Caulobacteraceae</i> (Proteobacteria)	15.7%	3.6%	0.0%	0.1%	0.0%	0.0%
<i>Xanthomonadaceae</i> (Proteobacteria)	28.6%	0.0%	0.4%	0.4%	4.2%	0.0%
<i>Xanthobacteraceae</i> (Proteobacteria)	1.0%	3.0%	4.0%	1.3%	0.8%	2.2%
<i>Comamonadaceae</i> (Proteobacteria)	26.1%	16.3%	5.8%	3.6%	8.3%	2.2%
<i>Hyphomicrobiaceae</i> (Proteobacteria)	0.0%	6.6%	0.7%	0.2%	0.8%	0.0%
<i>Beijerinckiaceae</i> (Proteobacteria)	2.9%	0.6%	0.0%	0.1%	0.0%	0.0%
<i>Methylophilaceae</i> (Proteobacteria)	0.1%	0.0%	0.7%	0.0%	2.5%	0.0%
<i>Rhodobacteraceae</i> (Proteobacteria)	0.0%	0.0%	1.4%	0.0%	4.2%	0.0%
<i>Burkholderiaceae</i> (Proteobacteria)	0.4%	2.4%	0.4%	0.0%	0.0%	1.1%
<i>Sutterellaceae</i> (Proteobacteria)	0.0%	0.0%	0.0%	0.1%	3.3%	1.1%
<i>Solimonadaceae</i> (Proteobacteria)	0.1%	0.0%	0.0%	0.0%	0.0%	3.3%
<i>Rhizobiales Incertae Sedis</i> (Proteobacteria)	0.0%	0.0%	2.2%	0.4%	0.8%	3.3%
<i>Gemmataceae</i> (Planctomycetota)	0.0%	1.2%	5.0%	1.5%	0.0%	2.2%
<i>Phycisphaeraceae</i> (Planctomycetota)	0.0%	7.8%	0.0%	0.2%	0.0%	0.0%
<i>Nitrospiraceae</i> (Nitrospirota)	0.0%	2.4%	3.2%	1.0%	1.7%	4.3%
<i>Chitinophagaceae</i> (Bacteroidota)	0.0%	0.0%	4.3%	1.4%	1.7%	1.1%
<i>Methylomirabilaceae</i> (Methylomirabilota)	0.0%	0.0%	0.0%	0.1%	0.0%	3.3%
<i>Planococcaceae</i> (Firmicutes)	0.0%	0.0%	0.4%	0.8%	0.0%	2.2%
<i>Bacillaceae</i> (Firmicutes)	0.0%	0.0%	0.7%	1.4%	0.0%	2.2%
<i>Sumerlaeaceae</i> (Sumerlaeota)	0.0%	0.0%	0.0%	0.2%	0.0%	2.2%
<i>Vicinamibacteraceae</i> (Acidobacteriota)	0.0%	0.0%	2.2%	0.7%	0.8%	0.0%
<i>Zixibacteria</i> (Zixibacteria)	0.0%	2.4%	0.0%	0.0%	0.8%	0.0%

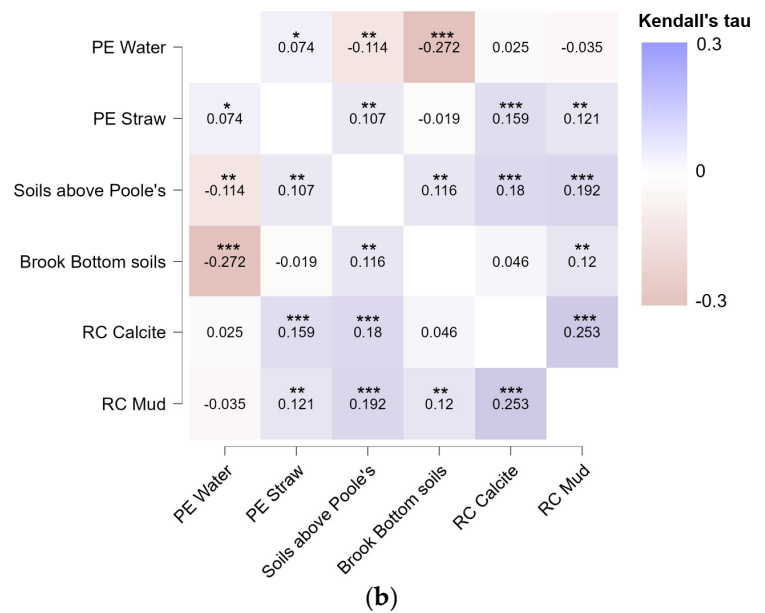
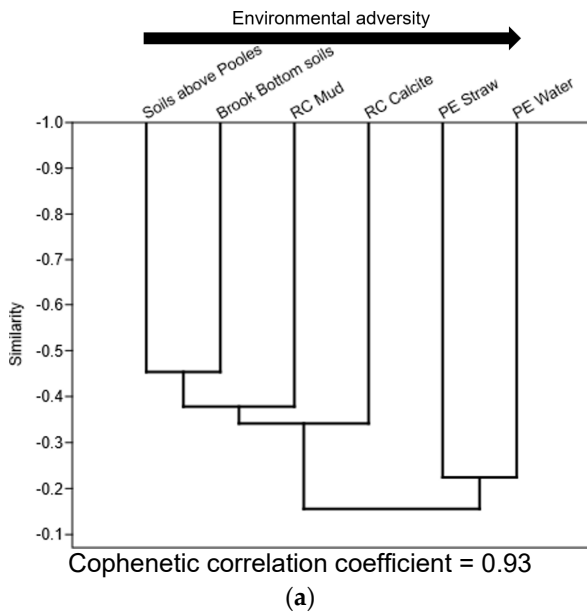
371 *Note.* Nomenclatures follow updated Silva 138 database.

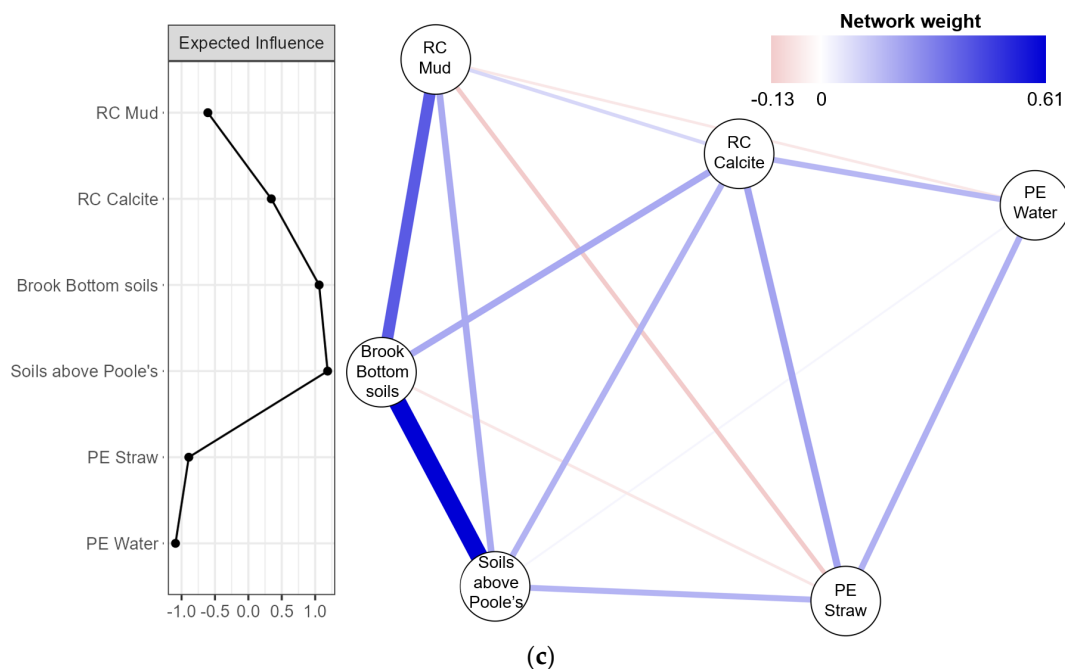
372 3.4 Ordination analyses of study sites

373 Based on the hierarchical clustering analysis of the sequence data, soils above Poole’s
 374 Cavern were the most similar to Brook Bottom soils (~45% similarity) (Figure 3a). Their cluster
 375 was the most similar to RC Mud (~38% similarity), and then to RC Calcite (~32% similarity).
 376 PE Straw and PE Water were clustered together but with the lowest level of similarity (~22%
 377 similarity).

378 Since none of the relative abundances of microbiomes from sampling sites demonstrate a
 379 normal distribution pattern, Kendall’s tau correlation coefficients were calculated and depicted in
 380 Figure 3b. Intriguingly, the PE Water sample significantly anticorrelated with the two soil
 381 samples – Soils above Poole’s and Brook Bottom soils. This trend indicated that most of the
 382 dominant microbial members in the least adverse soil samples outside Poole’s Cavern
 383 diminished in the harsh hyperalkaline water, while most of the rare species in external soils
 384 became dominant in the in-cave hyperalkaline water. Besides, the microbial community of PE
 385 Water was moderately associated with PE Straw, while PE Straw was highly associated with RC
 386 Calcite, and RC Calcite was highly associated with RC Mud. Additionally, the two external soil
 387 samples were correlated with each other as well as the samples from the RC chamber.

388 Moreover, network analysis that visualises the co-occurrence between species
 389 demonstrated that PE Water were the most marginal point that had the weakest connection with
 390 other sites; PE Straw was slightly less marginalized; two RC samples Mud and Calcite were
 391 more centralized. Centrality plot (represented by the expected influence index) delineated that
 392 Soils above Poole’s was the most influential site that was positively linked with other sites,
 393 followed by Brook Bottom soils (Figure 3c).





394 **Figure 3.** (a) Hierarchical clustering of Poole's Cavern samples using Bray-Curtis dissimilarity
 395 index based on bacterial species; (b) Kendall's tau correlation heatmap (*, $p < 0.05$; **, $p < 0.01$;
 396 ***, $p < 0.001$); (c) centrality and network plots.

397 By closely inspecting microbial classes, in-cave niches were distinct from Brook Bottom
 398 soils primarily due to abundance differences in Acidobacteriae, Holophagae, Vicinamibacteria,
 399 Acidimicrobiia, Thermoleophilia, Bacteroidia, Anaerolineae, Desulfuromonadia, Syntrophia,
 400 Bacilli, Clostridia, Latescibacteria, MBNT15, Methylomirabilia, Myxococcia, Polyangia,
 401 Planctomycetes, Alphaproteobacteria and Gammaproteobacteria. On the other hand, in-cave
 402 niches were distinct from Soils above Poole's primarily due to Vicinamibacteria, Acidimicrobiia,
 403 Bacteroidia, Polyangia, Nitrospiria, Phycisphaerae, Alphaproteobacteria and
 404 Gammaproteobacteria.

405 3.5 Metabolic pathway analyses

406 Of the different metabolic pathways identified in the Poole's Cavern samples, half were
 407 common across all samples and most involved the biosynthesis of essential biomolecules (Figure
 408 4). The most common of these metabolic pathways were: amine and polyamine biosynthesis,
 409 amino acid biosynthesis, aromatic compound biosynthesis, carboxylate degradation, cell
 410 structure biosynthesis, cofactor and vitamin biosynthesis, fatty acid and lipid biosynthesis, fatty
 411 acid and lipid degradation, nucleoside and nucleotide biosynthesis, nucleoside and nucleotide
 412 degradation, secondary metabolite biosynthesis, secondary metabolite degradation, aminoacyl-
 413 tRNA charging, fermentation, nucleic acid processing, pentose phosphate pathways,
 414 photosynthesis, respiration and tricarboxylic acid (TCA) cycle.

415 PE Water, as the sole liquid sample, showed higher proportions of aldehyde degradation,
 416 amino acid degradation and aromatic compound degradation, but lower relative proportions of
 417 alcohol degradation, carbohydrate biosynthesis, polymeric compound degradation, C1 compound
 418 utilisation and assimilation, electron transfer and glycolysis (Figure 4). PE Straw had relatively

419 lower proportions of alcohol degradation, aldehyde degradation and detoxification/antibiotic
 420 resistance than other sites. Soils above Poole's had a higher proportion of cofactor, prosthetic
 421 group and electron carrier degradation. Brook Bottom soils had higher proportions of alcohol
 422 degradation and Entner-Duodoroff pathways. RC Calcite had a higher proportion of amine and
 423 polyamine degradation, while RC Mud had a lower proportion of glycan degradation.



424
 425 **Figure 4.** Proportions of predicted metabolic pathway superclasses by PICRUSt2 in reference to
 426 the MetaCyc metabolism database.

427 The percentages of salinity-related pathways ranged from 1.03% in PE Water to 1.21% in
 428 RC Mud. The percentages of inorganic nutrient metabolism ranged from 1.45% in Brook Bottom
 429 soils to 1.76% in PE Straw (Table 4). Some pathways relevant to nitrate, sulphate and phosphate
 430 metabolisms contain assimilatory nitrate reduction, denitrification, nitrifier denitrification,
 431 assimilatory sulphate reduction, sulphur oxidation and methylphosphonate degradation.

432 **Table 4**

433 *Percentages of Predicted Pathways Related to Salinity and Inorganic Nutrient Metabolisms.*

Site	Metabolism	Salinity	Inorganic nutrient metabolism
PE Water		1.03%	1.61%
PE Straw		1.16%	1.76%
Soils above Poole's		1.17%	1.58%
Brook Bottom soils		1.20%	1.45%
RC Calcite		1.14%	1.70%
RC Mud		1.21%	1.51%

434 4 Discussion

435 Cave deposits, including speleothems, are prominently influenced by factors external to
436 the cave system. Inorganic minerals and organic molecules are released from the soil, aquifer and
437 bedrock and flushed into the cave system, where they are re-deposited on speleothem surfaces.
438 Therefore the “source” region is often defined as the soils and waters above the cave and the
439 vector of transport the percolating waters that eventually enter the cave. Whilst clear connections
440 can often be seen between physical events (large rainfalls or autumnal flushes) with influxes of
441 inorganic nutrients (Borsato et al., 2007; Wynn et al., 2014) and organic carbon (Baker et al.,
442 1993; Webb et al., 2014), less is known on how this continuous transfer of nutrients and minerals
443 and indeed microbial cells will influence the makeup of the in-cave microbial network (Zhu et
444 al., 2019). Additionally, the delivery of nutrients driven by environmental alterations such as UV
445 fluorescence laminations can vary notably on a secular basis (Webb et al., 2014) or an annual
446 basis (Baker et al., 1993). Understanding the individual community structures in the key source
447 environments (soils, waters) and those in the sinks (speleothems, muds) can help elucidate the
448 extent to which in-cave communities are a function of their source communities, or if they can
449 successfully adapt and diversify forming microhabitats or niches (Zhu et al., 2021).

450 4.1 Microbial community structures in different microhabitats

451 Proteobacteria represent the dominant phyla in all the samples collected within and
452 around Poole's Cavern. Most sites and sample types have a high abundance (if not dominant) of
453 previously discovered *Comamonadaceae* (Table 3), which is unsurprising as they are a versatile
454 bacterial family, including anaerobic denitrifiers, hydrogen oxidisers, ferric iron reducers,
455 photoautotrophs, photoheterotrophs and aerobic organotrophs (Burke et al., 2012; Willems,
456 2014). Apart from the dominance of Proteobacteria in all sites, the microbial community
457 structure from soils, waters, mud and speleothems in this hyperalkaline cave environment
458 (Hartland et al., 2010b) varied remarkably (Figure 2 & Table 2). As a rule, the biodiversity and
459 abundance of bacteria in karst ecosystems were less than soils above the cavern, indicating a
460 stronger selection pressure at the community level within the cave (Bassil et al., 2020).

461 4.1.1 Soils

462 The soil systems above Poole's Cavern are by far the most diverse in terms of community
463 structure, due to the organic rich and more neutral nature of the soils and the lack of requirement
464 for highly specialised microbial communities to develop. Only the Soils above Poole's Cavern
465 were characterised by *Gemmataceae*, *Chitinophagaceae* and *Vicinamibacteraceae* (Table 3). The
466 bacterial families found here, but not elsewhere, suggests a community well adapted to the
467 temperate forest ecosystem overlying the cave and include *Gemmataceae* which are aerobic
468 chemoorganotrophic bacteria that inhabit soils, wetlands, and freshwater (Kulichevskaya et al.,

469 2020); *Chitinophagaceae* which degrade chitin or even hydrolyse cellulose (Rosenberg, 2014),
470 and *Vicinamibacteraceae* which are neutrophilic and psychrotolerant chemoheterotrophs (Huber
471 and Overmann, 2018). The dominance of these specific families within the soil means they can
472 be used as tracers for “downstream” environments, helping us to identify if these communities
473 can be flushed through the soil system and into the cave, and once washed in if they can still
474 function/thrive in this unique environment of high pH waters and complete darkness.

475 4.1.2 Cave Waters

476 PE Water was the least diverse of all the samples collected, immediately indicating a
477 disconnect between the highly diverse soils above the cave and the water emanating from the
478 karst aquifer. Microbes in this water sample with pH>12 were even plausibly not metabolically
479 active (Smith et al., 2016), however were still viable as demonstrated by the presence of
480 culturable microorganisms. Only four families, *Sphingomonadaceae*, *Xanthomonadaceae*,
481 *Comamonadaceae* and *Caulobacteraceae* make up over 80% of the bacteria found within this
482 water (Table 3). The sole bacterial family that dominates PE Water only were *Beijerinckiaceae*
483 (Table 3), a group of nitrogen fixers and methanotrophs (Morawe et al., 2017). Whilst we are
484 limited to assessing a single sample, there was also no evidence for the potential tracer families
485 unique to the Soil above Poole’s (*Gemmataceae*, *Chitinophagaceae* and *Vicinamibacteraceae*)
486 within the drip water. This disconnect is clear in Figure 3a and 3b where there is no clear linkage
487 between the soils above Poole’s Cavern and the drip water emanating in PE. This, and the
488 reduction of culturable microorganisms in the water sample, indicates that leaching of microbial
489 communities from the soil to the aquifer is limited or that during transit through the karst,
490 significant community filtering/ restructuring occurs. In the drip waters two of the most
491 dominant families (*Sphingomonadaceae* and *Xanthomonadaceae*) are known to degrade
492 aromatic compounds (Balkwill et al., 2003) and reactive oxygen species (Saddler and Bradbury,
493 2005), quite different from the dominant soil bacteria. This reduction in the diversity of
494 microorganisms, and change in metabolic pathway, all indicate a significant reduction in suitable
495 micronutrients and environmental conditions to support a diverse microbial community within
496 the karst aquifer and resultant drip waters. These are significant observations when considering
497 the karst aquifer as a potential filter or zone of consumption for enhanced nutrient leachates from
498 anthropogenic activities.

499 4.1.3 Speleothem calcite

500 As mentioned in the sample collection section, the stalagmite samples from Poole’s
501 Cavern (PE High and Low) were grown on glass plates (watch glasses) and collected over a
502 relatively short time period (two months). These samples gave an opportunity to assess the rates
503 of biodiversification and the extent to which these “new” calcite samples varied from, for
504 example, the straw speleothem from the same chamber. Unfortunately, the PE High and Low
505 calcite samples showed the lowest extractable DNA abundance (Table 2) and there was not
506 enough extracted DNA to perform taxonomic phylogeny assessments. This lower abundance
507 could be due to a limited time for community development (probably due to rapid drip washing
508 rates) or could be a function (such as partially metabolic inhibition) (Smith et al., 2016) of a high
509 pH feeding water (pH 12.2), far more alkaline than Roman Chamber (pH 9.3) (Table 2).

510 Where we could perform an assessment of the dominant bacterial families within the
511 cave system, including PE Straw and RC Calcite, we found differences in the bacterial makeup

512 of these sampling sites. *Rhodobacteraceae*, *Sutterellaceae*, and *Methylophilaceae* were
513 characteristic families in the RC Calcite sample. *Rhodobacteraceae* are a universal but less
514 dominant marine family (Pohlner et al., 2019). *Sutterellaceae* are asaccharolytic anaerobic to
515 microaerophilic bacteria (Morotomi, 2014). *Methylophilaceae* are oligotrophs that consume
516 simple organics such as methanol or methylamine through dehydrogenases. The genomic
517 evolution of *Methylophilaceae* to adapt to environmental changes is primarily driven by
518 horizontal gene transfer (Salcher et al., 2019). These bacteria demonstrate essential traits to fit in
519 the oxygen- and nutrient-deficient in-cave conditions.

520 In comparison, *Caulobacteraceae* were enriched in PE Straw (3.6%) and PE Water
521 (15.7%); they are motile, with the presence of prosthecae and flagella, and alkaline phosphatase
522 is ubiquitous within this family to adapt to the high pH environment (Abraham et al., 2014). This
523 co-dominance indicates some interconnectivity between the drip waters and the speleothem
524 calcite developing from them and a clear adaption to the hyperalkaline environment in which
525 they survive. *Zixibacteria*, *Hyphomicrobiaceae*, *Phycisphaeraceae* and *Burkholderiaceae* were
526 dominant bacterial families only in the PE Straw sample. *Zixibacteria* have previously been
527 detected in moonmilk cave deposits (Maciejewska et al., 2018). *Hyphomicrobiaceae* can perform
528 denitrification or mixed-acid fermentation under anaerobic conditions (Oren and Xu, 2014).
529 *Burkholderiaceae* are saprophytic microbes (Coenye, 2014). *Phycisphaeraceae* are nitrate
530 reducing species discovered in marine algae that have alkaline phosphatase (Fukunaga et al.,
531 2009). Metabolic pathways of inorganic nutrient metabolism within cave environments hold
532 important status for understanding the nutrient content of speleothem calcite (Wynn et al., 2021;
533 Wynn et al., 2008). The finding of these microbial inhabitants of the straw but not in the PE
534 Water is significant, implying that this community has developed independently within the cave,
535 and not controlled by the water source feeding the precipitating calcite.

536 Due to the nature of the PE Straw sample this may only reflect part of the potential
537 community in these calcite drip samples. Although some researchers have found minimal variety
538 within calcite samples in a cave system (Dhami et al., 2018), more of them have found reduced
539 similarity (Mendoza et al., 2016; Ortiz et al., 2013; Park et al., 2020; Van de Kamp, 2004).
540 Despite this ambiguity the differences in the communities between PE Straw and PE Water
541 reflect a disconnect between the water and some areas of the cave system.

542 4.1.4 Sediments/ mud

543 The RC Mud sample has a similar level of diversity as the RC Calcite collected from the
544 same chamber, with *Xanthobacteraceae* (2.2%) and *Nitrospiraceae* (6.7%) slightly dominant in
545 RC Mud, similar to Soils above Poole's. However, RC Mud had the most bacterial families
546 many of which were not found in other in-cave samples (Table 3), i.e., *Solimonadaceae*,
547 *Methylomirabilaceae*, *Planococcaceae*, *Bacillaceae* and *Sumerlaeaceae*. One of the dominant
548 families, *Nitrospiraceae*, are nitrite oxidisers (Daims, 2014) capable of metabolising under
549 hyperalkaline conditions (Daebeler et al., 2020), reflective of the conditions even in the RC
550 sediments which are fed by hyperalkaline waters. The family *Solimonadaceae* is primarily found
551 in soils and freshwater; many species of *Solimonadaceae* can decompose chemical pollutants
552 such as chlorinated compounds (Zhou et al., 2014). *Methylomirabilaceae* are denitrifying
553 methanotrophs that couple anaerobic methane oxidation with nitrite reduction to nitrogen (Ettwig
554 et al., 2010). *Planococcaceae* are more abundant in terrestrial habitats rather than aquatic or
555 marine environments (Gupta and Patel, 2020). *Bacillaceae* are endospore-forming bacteria (Vos

556 et al., 2011). *Sumerlaeaceae* are rarely reported but they have been identified in other extreme
557 environments, including cold arid deserts and deep-sea basins (Fang et al., 2021). As above,
558 these families can all adapt to soils or moister environments. More dominant families suggest
559 that the less alkaline mud below speleothems can foster a more diverse community.

560 The dominant bacterial families of Poole's Cavern microbiomes were not different from
561 most cave and karst systems (Engel, 2007, 2010; Farnleitner et al., 2005; Ikner et al., 2007;
562 Northup et al., 2003; Pronk et al., 2009), although far fewer common families are identified
563 (Bassil et al., 2015). These findings highlight that there are significant adaptations found within
564 the cavern in comparison to the soils or waters feeding the in-cave locations. Disconnections
565 between sites suggest that the microbial communities in the soil-karst-speleothem-sediment
566 system are not part of a continuum, but only a partial continuum, with opportunity for adaptation
567 and unique community development specific to the immediate environmental conditions and
568 substrates on which they are supported.

569 4.2 Microbial network between sites

570 As discussed above, alkaliphilic microbial dwellers of Poole's Cavern samples possessed
571 various metabolic functions to consume simple inorganic or organic nutrients and to adapt to
572 changes in ions (Table 4). The main bacteria taxa in Poole's Cavern are similar to neutral to
573 slightly alkaline cave environments (Hershey and Barton, 2018; Tomczyk-Zak and
574 Zielenkiewicz, 2016). Some bacterial taxa were common between sites, and some subgroups
575 such as *Methylophilaceae* develop new metabolic functions by horizontal gene transfer (possibly
576 to adapt to the different environments of RC Mud and RC Calcite, Table 4 & Figure 3b, c).
577 Hence, the network between microbial communities from different microhabitats can be crucial
578 in a complex cave system. The anticorrelation between PE Water and two external soil niches
579 implied that dominant bacterial inhabitants in milder environments lost their advantages in a
580 hyperalkaline aquatic setting (Figure 3b). Additionally, proximity of PE Straw to the Soils above
581 Poole's in the correlation and network graphs suggest a rapid hydrological connection from the
582 surface (Figure 3b, c), implying less time for species specialisation. Despite the PE Straw being
583 more hyperalkaline at the drip site, the dry surface conditions of the straw feed the microbes in a
584 similar manner to RC Calcite regarding microbial compositions (Figure 3b).

585 However, whilst commonality of bacterial taxa between sites was present, and functional
586 likeness was evidenced in specific circumstances, this connectivity was weak (network weight <
587 0.21) among in-cave sites and the change in ecological niches can lead to shifts in microbial
588 consortia (Park et al., 2020). Slightly different from previous microbiological studies of less
589 alkaline caves (Dhami et al., 2018; Hershey and Barton, 2018; Tomczyk-Zak and Zielenkiewicz,
590 2016), both the hierarchical clustering (similarity < 45%) and network analyses displayed
591 relatively lower similarity among different sampling sites (Kruskal-Wallis H of inter-site
592 difference = 443.8, $p < 1.1 \times 10^{-93}$), indicating that microbial communities of these sites interacted
593 and affected each other less often (Figure 2 and Figure 3). Despite reduced microbiota
594 connections, the positive associations between RC Calcite and RC Mud, as well as PE Water and
595 PE Straw manifested the presence of a spatial continuum within Poole's Cavern chambers
596 (Figure 3b), as Blyth et al. (2014) also proposed that organic molecules such as biolipids in cave
597 speleothems and associated soils were likely derived indigenously (Blyth et al., 2014).

598 Since the connectivity between microhabitats in Poole's Cavern was not high, each
599 microbial group from its respective ecological niche has to rely on its own metabolisms. Within

600 this soil-water-speleothem-sediment continuum, microbes evolve and adapt to their respective
601 microhabitats along the environmental adversity gradient (Figure 3a), gradually forming distinct
602 communities, possibly also affected by their unique external sources that others are not exposed
603 to. Due to the changes in pH and nutrients, multiple functions were constructed *in situ*. Complete
604 independent metabolic function structure from each sampling site was detected using the
605 PICRUSt2 metagenome inference method (Figure 4). Almost all sites shared similar proportions
606 of essential biosynthesis pathways for producing amino acids, nucleotides, fatty acids, cell
607 structure materials, and amines. Other pathways varied among sites to adapt to their unique
608 environmental conditions. Microorganisms in each site were capable of utilising various and
609 sometimes even unusual inorganic or organic compounds to survive (Charles et al., 2015). The
610 dominance of pathways is closely related to environmental properties. The higher proportions of
611 salinity-related response pathways in mud samples than in the water sample (Table 4) were
612 probably due to evaporation during spring/summer. PE Straw and RC Calcite samples were less
613 trophic than the soils (Dhami et al., 2018), and thus had more percentages of inorganic molecule
614 consuming pathways. The evolution rates of microbial communities are quick in these
615 susceptible microhabitats, and bacteria are very versatile to develop strategies to survive in the
616 Poole's Cavern system.

617 **5 Conclusions**

618 Poole's Cavern in the UK is famous for its hyperalkaline speleothem deposits and
619 anthropogenically impacted karst formation history. Extremophilic microorganisms were
620 considered to dominate this region. In this study, the abundance, taxonomy and functions of
621 microbial communities from four types of samples (soil, water, speleothem and mud) were
622 examined to understand the distribution of microbial communities and their connections. The
623 aqueous conditions in Roman Chamber, located close to the cave entrance, were warmer, less
624 saline, and less alkaline than Poached Egg Chamber. Consequently, Roman Chamber nurtured
625 more abundant and diverse microbial communities. We discovered limited similarity between
626 different niches with community structure demonstrating a high variability among sampling
627 sites, feasibly driven by the steep alkalinity gradient. Each site possessed a relatively
628 independent complete essential metabolic structure due to the demands of self-sustainability.
629 This is the first study to identify microbial taxa that can adapt to an alkaline-hyperalkaline cave
630 ecosystem and inhabit different niches of a cavern and provide a basis to determine the controls
631 on these niches and potential seasonal variability. Investigations of alkaliphilic microorganisms
632 contributed to the understanding of global biosphere and microbe-environment interactions in
633 karst and cave science.

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642 **Open Research**

643 The 16S rRNA gene sequences used for taxonomic analysis and functional pathway
644 inference in this study are available in the NCBI BioProject database via the accession number
645 PRJNA807843 [BGS © UKRI]. A supplemental table Table S1 is available at the link:
646 <https://doi.org/10.6084/m9.figshare.20113907.v1>.

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