



1 **Geochemical and microbial factors driving crustacean assemblages** 2 **in adjacent groundwater bodies within the same aquifer**

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19 **Abstract.** Aquifers harbor unique and highly adapted species, contributing to critical ecological processes and services.
20 Understanding the key factors driving invertebrate assemblages in aquifers is a challenging task that, traditionally, has
21 primarily been achieved in karst aquifers. This study aimed to uncover the factors influencing the composition and
22 functionality of groundwater crustaceans in a volcanic aquifer of central Italy. We adopted a multidisciplinary approach,
23 combining hydrogeology, geology, microbiology, and ecology, and found that the aquifer consisted of three adjacent
24 groundwater bodies (GWBs) with different geochemistry (i.e., sulfate-depleted, K-rich and earth-alkaline) and microbial
25 characteristics that remained consistent over the study period. We also unveiled significant differences in both the taxonomic
26 and functional composition of groundwater crustaceans across the three GWBs and these patterns were consistent over time.
27 Notably, the sulfate-depleted GWB lacked groundwater-obligate species, burrowers, stenothermal and moderately
28 stenothermal species, while the K-rich and earth-alkaline GWBs had different species with similar functions related to
29 locomotion, diet, and feeding habits. Stenothermal and moderately stenothermal crustacean species were only found in the
30 K-rich GWB, which lacked epigeal species. Major ions (SO₄, Ca, NO₃, and K), trace elements (B, Al, V, Se, and Ba), and
31 microbial factors related to microbial cells with low nucleic acid (LNA cells) and carbohydrate catabolic profiles were the
32 main descriptors of groundwater-obligate species abundances. Our findings revealed a significant correlation between the
33 abundances of groundwater-obligate crustaceans and LNA cells, thus suggesting a selective feeding of groundwater
34 invertebrate species on the aquatic microbial community. Our research emphasizes the need to consider diverse
35 hydrogeological contexts within individual aquifers. Potential avenues for future research should further consider food web
36 dynamics in groundwater communities and their impact on carbon and nutrient cycling.



37 **1 Introduction**

38

39 The Earth's largest unfrozen reserve of freshwater is found within aquifers, geological formations responsible for storing and
40 transmitting groundwater (Ferguson et al., 2021). According to geological and hydrological criteria, aquifers can be divided
41 into specific sections known as groundwater bodies (GWBs) (Aquilina et al., 2023). They harbor highly specialized
42 microbial and metazoan species adapted to life in permanent darkness and low energy conditions (Malard et al., 2023a).
43 Groundwater microbes and metazoans play crucial roles in providing ecosystem services such as carbon recycling, pathogen
44 removal, pollutant bioremediation, sediment mixing, and burrowing (Fillinger et al., 2023; Mermillod-Blondin et al., 2023).
45 The variation in the taxonomic and functional composition of the invertebrate assemblages of a given aquifer across its
46 groundwater bodies results from a complex interplay of abiotic (e.g., hydrochemistry, porosity, flow velocity) and biotic
47 factors (e.g., species competition, cross-kingdom interactions), acting on both evolutionary and ecological scales (Zagmajster
48 et al., 2023). However, disentangling the combined effects of these taxonomic and functional drivers remains a difficult task
49 (Malard et al., 2023b).

50 The species' ability to spread across different groundwater bodies depends on extrinsic factors (e.g., the degree of
51 connectivity among adjacent water bodies) and intrinsic characteristics of the species, which are mainly defined by the
52 suitability of its functional traits (e.g., body size, shape, locomotion, and feeding habits) and the ability to adjust or adapt its
53 traits in response to the new habitat template (Fišer et al., 2023). If a species lacks trait plasticity, it is likely to struggle to
54 establish itself in a new environment. This phenomenon is known as habitat (or environmental) filtering (Cornwell, Schilck,
55 and Ackerly, 2006; Kraft et al., 2015).

56 Groundwater crustaceans, which are the dominant metazoans in global groundwater ecosystems (Stoch and Galassi, 2010),
57 are known to undergo environmental filtering due to abiotic factors such as pore size, flow rate, connectivity to the surface,
58 and hydrochemistry (Trontelj, Blejec, and Fišer, 2012). The role of trophic filters has been less investigated. Groundwater
59 crustacean species in a given aquifer compete for space and food resources (Bregović et al., 2019). Traditionally, research on
60 the effect of environmental filtering on groundwater crustaceans has primarily focused on karst aquifers (Culver et al., 2021).
61 However, there is a growing recognition of the importance of non-karst aquifers, such as the volcanic ones (Wurzbacher et
62 al., 2020), which has prompted an intensified effort to gain a better understanding of groundwater ecosystems across a wide
63 range of hydrogeological contexts (Hahn and Fuchs, 2009).

64 In this study, we aimed to unravel the factors shaping groundwater crustacean assemblages within a water table aquifer
65 where groundwater circulates in volcanic and sedimentary formations. Preliminary analyses carried out two years prior to
66 this study indicated the occurrence of adjacent, chemically different, GWBs within the same aquifer (Ademollo et al., 2012).
67 Accordingly, the objective of our study was to investigate potential disparities in the taxonomic and functional composition
68 of the crustacean assemblages along with the spatial variations of the aquifer hydrochemistry. More specifically, our goal
69 was to pinpoint the primary factors that influence the composition of crustacean assemblages across the three GWBs,
70 including: i) hydrogeological factors (e.g., water table depth, permeability and porosity of the rock matrix); ii) physical-

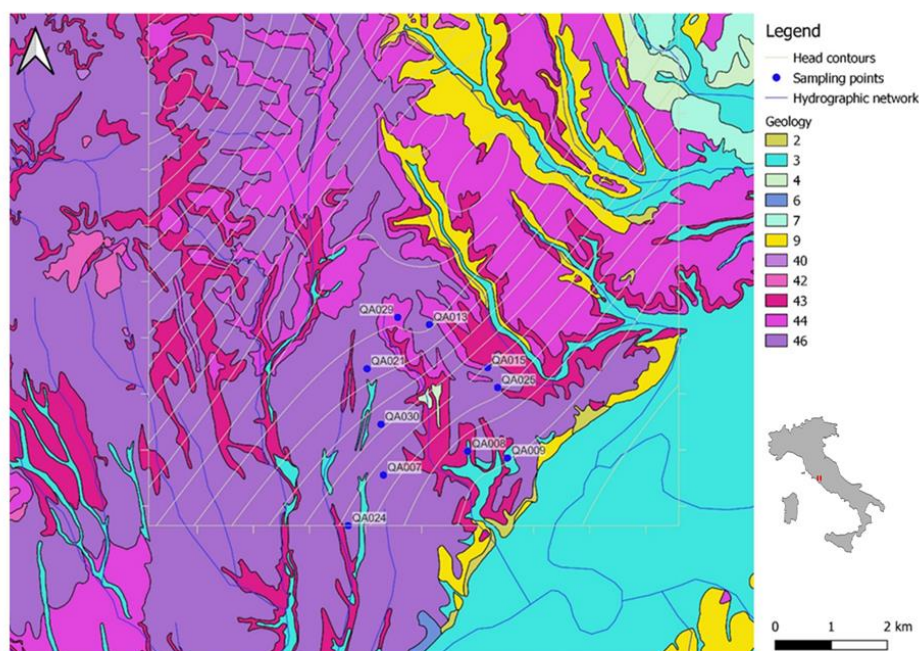


71 chemical factors encompassing temperature, pH, dissolved oxygen, as well as major ions and trace elements; iii) microbial
72 factors, such as microbial cell abundance and metabolic profiles. This study ventures into relatively uncharted territory of
73 volcanic aquifers and addresses critical gaps in our understanding of the main factors shaping the groundwater invertebrate
74 assemblages.

75 2 Materials and Methods

76 2.1 Hydrogeological aquifer settings and sampling survey

77 We conducted our study on the eastern flank of the Sabatini Mounts aquifer, near the valley of River Tiber, covering an area
78 of 30 square kilometers in the northeastern province of Rome (Italy). The sampling site is in a rural area across the Fosso
79 Fontanalarga, a minor right-hand tributary of the River Tiber, with small-medium villages, sparse houses, agricultural fields
80 and a few industrial activities, including a quarry for the exploitation of lapideous volcanic products (namely the “Via
81 Tiberina Yellow Tuff” formation; Lombardi and Meucci 2006). Pleistocene K-alkaline volcanic products of the Roman co-
82 magmatic province (Sottili et al., 2010), mainly pyroclastic such as ignimbrites and tuffs, overlap in angular unconformity
83 Pliocene and Pleistocene sedimentary deposits of marine and continental origin (Fig. 1).



100 **Figure 1: Geological map of the study area and sampling point locations. Head contours referring to 2014 survey are plotted.**
101 **Geology: 2) colluvial-alluvial covers (Olocene); 3) Alluvial deposits (Holocene); 4) Lacustrine-marsh deposits (Holocene); 6)**
102 **Ancient alluvial terraces deposits (Pleistocene); 7) Travertines (Pleistocene); 9) Marine sands and gravels (Pleistocene); 40) tephra;**
103 **42) Trachitic lavas (Pleistocene) 43) Lithoid ignimbrite (“Via Tiberina Yellow Tuff”).**



104 Holocene alluvial deposits can be found in the valley talwegs. The water table aquifer is hosted in the volcanites, with
105 groundwater circulating mostly in the porous and fractured lithoid levels of the ignimbrites and lavas, and in the marine and
106 continental sands and gravels below. Low permeability levels of the Pliocene marine clays and silts form the bottom of the
107 aquifer and isolate it from the deep geothermal reservoir hosted in the thick sequence of the Meso-Cenozoic fractured
108 limestones below (Parrone et al., 2020). The Pleistocene pyroclastic products, referred to herein as volcanites, typically
109 exhibit a medium permeability. Since the volcanic formations outcrop more extensively (Fig. 1), aquifer recharge occurs
110 mainly from these rocks than from the underlying sedimentary layers. Groundwater, mainly exploited for irrigation and to
111 supply households not connected to the public drinking water supply system, flows at depth from north-west to south-east
112 direction through both volcanic and sedimentary layers, the latter being more important towards the edge of the volcanic
113 apparatus (Preziosi et al., 2013; Parrone et al., 2019).

114 **2.2 Field and laboratory geochemical analysis**

115 Preliminary data, collected in March 2012 (Ademollo et al., 2012), indicated the occurrence of three adjacent GWBs with
116 distinct hydrogeochemical features, characterized by sulfate-depleted, earth-alkaline, and K-rich waters, respectively. In this
117 study, we conducted two sampling campaigns: the first was carried out between November and December 2014, and the
118 second took place from October to December 2015. During these campaigns, we collected 20 groundwater samples from a
119 total of 10 wells (Fig. 1; Table A1). The distribution of the wells was as follows: i) wells QA7, QA21, QA24, and QA30,
120 situated on the western side of the aquifer, were associated with the GWB characterized by sulfate-depleted groundwaters; ii)
121 wells QA8, QA9, and QA29, situated along the valley of Fosso Fontanalarga, were within the GWB with earth-alkaline
122 groundwaters; iii) wells QA13, QA15, and QA25, located on the eastern side, belonged to the GWB characterized by K-rich
123 groundwaters. For each well, we recorded essential field data, including GPS coordinates, elevation and depth of the well,
124 depth to water table, oxidation-reduction potential (ORP), temperature, pH, dissolved oxygen, and electrical conductivity
125 using multiparametric portable probes (Aquaread AP 2000). To ensure accurate measurements, we purged the well until the
126 physical-chemical parameters stabilized before sampling. Water samples for chemical analyses were filtered on-site with
127 0.45- μm membrane filters and stored in HNO_3 1% treated polyethylene bottles. For major cations and trace metal
128 determination, samples were stabilized by adding HNO_3 at pH 2. Alkalinity was determined in the laboratory by HCl
129 titration on 50 mL of sample within 24 h from sampling. Anions were determined by ion chromatography (IC, Dionex DX-
130 120), major cations by ICP-OES (Perkin Elmer P400) and trace elements by inductively coupled plasma mass spectrometry
131 (ICP-MS, Agilent technologies 7500c). Samples for dissolved organic carbon (DOC) were filtered at 0.7 μm with pretreated
132 fiber glass filters and DOC levels were determined through a Shimadzu TOC-5000 analyzer. Further analytical details are
133 reported elsewhere (Preziosi et al., 2014; Parrone et al., 2022).

134

135 **2.3 Microbial and crustacean community analyses**



136 Samples for the microbial and crustacean community analyses were collected from all wells after purging. Aliquots (2 mL)
137 were fixed with formaldehyde (2% final concentration), stored at 4°C, and analysed within one week from sampling by the
138 Flow Cytometer A50-micro (Apogee Flow System, Hertfordshire, England), equipped with a solid-state laser set at 20 mV
139 and tuned to an excitation wavelength of 488 nm. Total cell counts (TCC) were quantified using side scatter and green
140 fluorescence measurements. The intensity of green fluorescence emitted by SYBR-positive cells facilitated the distinction of
141 cell groups based on their nucleic acid content, namely cells with Low and High Nucleic Acid content (LNA and HNA cells)
142 (Amalfitano et al., 2018).

143 Additional aliquots (15 ml) were collected, stored in sterile Falcon tubes and promptly incubated within a 6-hour timeframe
144 to evaluate the functional diversity and metabolic preferences of the bacterial communities by measuring their degradative
145 activity toward a set of organic carbon sources using Biolog™ EcoPlates assay (Biolog, Inc., Hayward, California, USA).
146 The assay consists of a 96-well microplate that contains 31 common carbon sources categorized into five compound groups
147 (i.e., carbohydrates, polymers, carboxylic acids, aminoacids, and amines), plus a control well, in triplicate. The provided
148 substrates are labeled with the respiration-sensitive tetrazolium dye, which is reduced to formazan when microbial
149 degradation occurs. This is a colorimetric reaction with the typical purple color development, which optical density (OD) is
150 detected at the wavelength 590 nm by a multi-plate reader (Perkin Elmer VICTOR™ X3). OD values were measured
151 immediately after sample inoculation (T0) and after 24 hours (T24) of incubation in dark conditions at 20°C. Microbial
152 metabolic fingerprinting was expressed as the percentage value of each single class of compounds compared to the total OD
153 (Melita et al. 2019; Preziosi et al., 2019).

154 Additionally, we conducted in-situ filtration of 1000 L of water from each well using a 63-µm mesh net to collect
155 groundwater invertebrates. The collected samples were preserved in a 70% ethanol solution and transported to the laboratory
156 for further analysis. In the laboratory, we sorted the samples using a stereomicroscope at 16× magnification, identifying the
157 taxa down to the species level, following established taxonomic references (e.g., Dussart and Defaye, 2006). Each specimen
158 was categorized as either groundwater-obligate species (which complete the whole life cycle in groundwater) and epigean
159 species (which occasionally occur in groundwater but lack adaptations for a permanent dwelling). Subsequently, we
160 evaluated four functional traits, each defined by two or more categories. The traits encompassed locomotion, diet, feeding
161 habits, and thermal tolerance. To obtain the trait profile of each sample, we calculated the percentage of each trait category
162 based on its abundance within the sample (Di Lorenzo et al., 2021).

163

164 **2.4 Statistical analyses**

165 To confirm the occurrence and stable nature of the three GWBs, as assumed from our preliminary investigation in 2012, we
166 performed permutational analyses of variances (PERMANOVA; Anderson, 2008) based on physical-chemical factors
167 measured *in situ* (water table depth, temperature, pH, dissolved oxygen and electrical conductivity), major ions and elements
168 in trace, respectively. We conducted two-way PERMANOVAs, incorporating a factor "groundwater body" with three levels



169 (sulfate-depleted, earth-alkaline and K-rich) and another factor "year" with two levels (2014, 2015), using resemblance
170 matrices based on Euclidean distances calculated from normalized data. To ensure the robustness of our analysis, we
171 examined potential multicollinearity among the variables using Draftsman's plots prior to PERMANOVAs. Variables
172 exhibiting high collinearity ($|r| \geq 0.95$ in correlation) were considered to convey essentially identical information and were
173 consequently removed to prevent redundancy in the analysis, in line with recommendations in Anderson et al. (2008). The
174 variables retained for analysis serve as proxies for those that were eliminated (Anderson et al., 2008). We excluded oxygen
175 saturation from the analyses due to its $> 99\%$ linear correlation with dissolved oxygen. In line with best practices, we applied
176 permutation of residuals under a reduced model and employed Type III sum of squares with 999 permutations. This
177 approach offers high statistical power and more accurate type I error control for multi-factorial, unbalanced designs
178 (Anderson et al., 2008). When appropriate, we conducted permutational post-hoc t-tests.

179 We evaluated the completeness of crustacean sampling effort by examining the accumulation of the total number of different
180 observed species (Sobs) as samples were progressively added (Magurran, 2021). To assess the potential increase in species
181 richness (S) with repeated sampling, we applied five non-parametric (Chao1, Chao2, Jackknife1, Jackknife2, and Bootstrap)
182 and one parametric (Michaelis-Menten) estimators (Magurran, 2021). We computed the estimators at each stage as new
183 samples were added, resulting in the generation of six curves illustrating the progression of the S with increasing sample
184 size. We conducted the analyses through 999 randomizations without replacement (Magurran and McGill, 2011).

185 To investigate potential differences in the taxonomic and functional composition of crustacean assemblage and microbial
186 community among the three GWBs, we applied the PERMANOVA design previously outlined for the environmental
187 variables. We $\log(x+1)$ -transformed crustacean abundances and microbial cell counts before generating Bray-Curtis
188 resemblance matrices, while percentage data pertaining to metazoan functional traits and HNA and LNA cells remained
189 untransformed (Anderson, 2008). We added a dummy variable equal to 1 to all data to allow the analysis of values equal to
190 zero (Clarke and Gorley, 2005). We chose not to analyze the abundances of each crustacean species due to the diminished
191 interpretive accuracy when the abundances of individual taxa constituted less than 4% of the total abundances (Clarke and
192 Gorley, 2005). In our study, only four species met or exceeded this specified threshold (Sect. 3.2). While PERMANOVA
193 inherently does not necessitate explicit assumptions regarding the distributions of the original variables, we opted to conduct
194 a Levene's test using the PERMDISP routine prior to all analyses. We focused on PERMANOVA outcomes that were not
195 influenced by bias due to variance heterogeneity (Anderson, 2008). To provide a comprehensive overview of the significant
196 outcomes derived from the PERMANOVAs, we utilized boxplot when considered insightful for visualization.

197 Finally, to assess the main hydrogeological, physical-chemical and microbial factors that influence the composition and
198 functionality of crustacean assemblages across the three GWBs, we employed distance-based linear models (DisTLM) based
199 on the Bray-Curtis resemblance matrix (Legendre and Anderson, 1999). We conducted both conditional and marginal tests.
200 Conditional tests involved fitting one factor after another. We applied the BEST procedure to construct models utilizing the
201 best factor combination. We assessed the AICc value (Akaike's Information Criterion corrected for small sample sizes;
202 Hurvich and Tsai, 1993) for all possible combinations of predictor variables, with the smallest AICc value indicating the



203 most suitable model. Additionally, we used R^2 to evaluate the proportion of explained variation in the multivariate models.
204 The significance of the marginal tests was determined by computing the p-values through permutations rather than
205 traditional tables (Legendre and Anderson, 1999). For each test, we employed 999 permutations to obtain p-values testing
206 the null hypothesis of no relationship, either for individual variables in isolation or in a conditional context (Legendre and
207 Anderson, 1999). Factors that individually (marginal models) or together with others (conditional models) explained > 65%
208 of the variance in the taxonomic and functional structure of the crustacean assemblage were considered robust, following the
209 criteria established by Korbel and Hose (2011, 2017) and Di Lorenzo et al. (2020). We applied this cut-off criterion to
210 prevent unreliable and exaggerated claims about scientific phenomena (Kimmel et al., 2023). We performed distance-based
211 redundancy analyses (dbRDA; Legendre and Anderson, 1999) to visualize the ordination of fitted values from the most
212 robust models.

213 Significance levels (α) were set at 0.05 for all permutational tests since they provide an exact test of each individual null
214 hypothesis of interest (Anderson et al., 2008). All analyses were performed using E-PRIMER version 6 and
215 PERMANOVA+ software (Anderson et al., 2008). Boxplots were generated using the libraries ggplot2 and gridExtra and the
216 R software (R Development Core Team, 2021).

217 **3 Results**

218 **3.1 Environmental factors and microbial community patterns**

219 Our analyses confirmed that the aquifer was composed of three adjacent GWBs, each distinguished by its unique
220 hydrochemical facies. The analyses revealed no significant differences in the water table depth among the three GWBs or
221 between the two years (Table A2a and S1). However, PERMANOVA uncovered differences in temperature, pH, and
222 electrical conductivity (Table A2a and S1). In detail, the sulfate-depleted GWB exhibited a mean temperature exceeding that
223 of the other two GWBs by about 1 °C (Table 1). The earth-alkaline GWB showed the lowest pH and the highest electrical
224 conductivity values (Table 1; Fig. 2). However, no discernible distinctions were observed between the two years (Table S1).
225 Despite belonging to the same aquifer and geological facies, the three GWBs exhibited significant differences in the major
226 chemical components, with variations attributed to SO_4 , HCO_3 , Na, K, and Si (Table A2b and S1; Fig. 2). In detail: the
227 earth-alkaline GWB displayed the highest HCO_3 and Si concentrations; the K-rich GWB showed the highest mean
228 concentration of Na and K; the sulfate-depleted GWB displayed the lowest mean values of SO_4 and HCO_3 (Tables 1 and S1;
229 Fig. 2). Significant distinctions between the two years were only observed for Si and DOC, both of which were higher in
230 2014 compared to 2015 (Tables A2b and S1). Significant differences among the three GWBs were also noted for trace
231 elements, namely in the concentrations of Li, B, Rb, and U (Tables A2c and S1). The K-rich GWB showed the highest
232 concentrations of Li, B, and Rb, while the sulfate-depleted GWB showed the lowest concentrations of U (Table 1 and Fig.
233 2). No significant differences emerged between the two years for trace elements.

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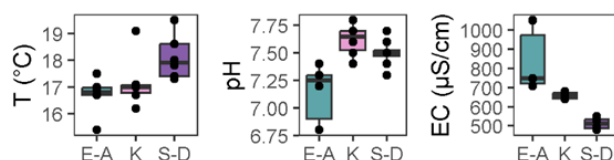
235 **Table 1: Physical-chemical and microbial characteristics of the groundwater samples in the three groundwater bodies (GWBs) of**
 236 **the target aquifer. Field parameters, major components, trace elements and microbial community properties are reported as**
 237 **means. Superscript letters (a, b, c) indicate statistical differences among GWBs (permutational post-hoc t-tests; $p < 0.05$).***
 238 **indicates statistical differences between 2014 and 2015.**

	Sulfate-depleted	Earth-alkaline	K-rich
239	Water table depth (m)		
240	80.01	64.26	81.82
241	Field parameters		
242	EC ($\mu\text{S}/\text{cm}$)	512.0 ^b	885.5 ^a
243	DO (mg/L)	7.3	5.9
244	DO (%)	77.3	61.2
245	ORP (mV)	183.7	156.6
246	pH	7.5 ^b	7.1 ^a
247	T ($^{\circ}\text{C}$)	18.1 ^b	16.8 ^a
248	Major components		
249	F (mg/L)	1.4	0.9
250	Ca (mg/L)	41.3	84.3
251	Cl (mg/L)	28.1	29.3
252	DOC (mg/L)	0.8	0.8
253	HCO ₃ (mg/L)	231.0 ^b	377.5 ^a
254	K (mg/L)	34.9 ^a	31.9 ^b
255	Mg (mg/L)	8.2	12.3
256	Na (mg/L)	24.6 ^a	31.9 ^{a,b}
257	NO ₃ (mg/L)	10.2	25.8
258	Si (mg/L)	22.7 ^a	22.8 ^{a,b}
259	SO ₄ (mg/L)	6.2 ^b	59.5 ^a
260	Trace elements		
261	Al ($\mu\text{g}/\text{L}$)	10.6	11.1
262	As ($\mu\text{g}/\text{L}$)	20.6	22.6
263	B ($\mu\text{g}/\text{L}$)	64.9 ^b	89.5 ^a
264	Ba ($\mu\text{g}/\text{L}$)	72.3	81.5
265	Cr ($\mu\text{g}/\text{L}$)	1.3	1.0
266	Cu ($\mu\text{g}/\text{L}$)	0.3	0.7
267	Fe ($\mu\text{g}/\text{L}$)	23.4	22.5
268	Li ($\mu\text{g}/\text{L}$)	18.6 ^b	15.5 ^a
269	Mn ($\mu\text{g}/\text{L}$)	0.9	1.9
270	Ni ($\mu\text{g}/\text{L}$)	0.3	0.7
271	Rb ($\mu\text{g}/\text{L}$)	21.7 ^b	29.9 ^a
272	Se ($\mu\text{g}/\text{L}$)	0.3	0.8
273	Sr ($\mu\text{g}/\text{L}$)	529.4	813.3
274	U ($\mu\text{g}/\text{L}$)	3.1 ^b	23.4 ^a
275	V ($\mu\text{g}/\text{L}$)	30.9	28.4
276	Zn ($\mu\text{g}/\text{L}$)	37.3	10.8
277	Microbial community		
278	TCC (10^4 cells/mL)	1.0 ^b	3.1 ^a
279	LNA cells (% of TCC)	65.7 ^b	75.6 ^a
280	HNA cells (% of TCC)	34.3 ^b	24.4 ^a
281	HNA/LNA	0.5 ^b	0.3 ^a
282	Carbohydrates	21.6	38.9
283	Polymers*	46.0	13.2
284	Carboxylic acids*	24.1	23.6
285	Aminoacids*	5.7	19.2
286	Amines	2.7	5.2

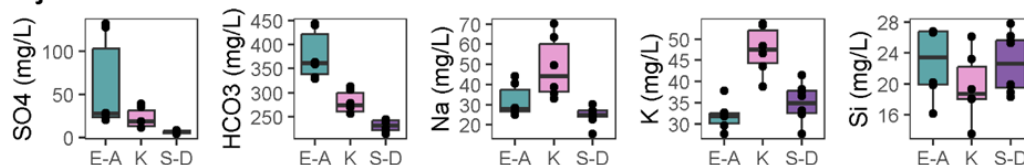


269 Total counts of microbial cells ranged from 0.5 to 9.5×10^4 cells/mL, with the lowest abundances occurring in the sulfate-
 270 depleted GWB (Table A2d and Fig. 2). A similar pattern was observed for the percentages of HNA and LNA cells, with the
 271 lowest percentages of LNA cells found in the sulfate-depleted GWB, which was correspondingly richer in HNA cells
 272 compared to the K-rich and earth-alkaline GWBs (Tables 1 and A2d; Fig. 2).
 273

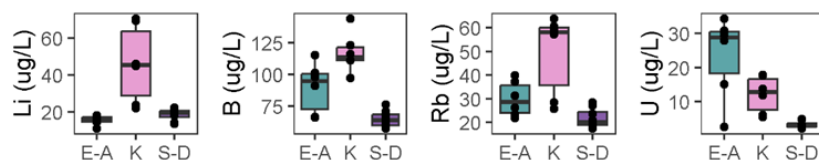
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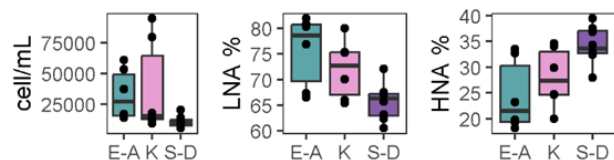
Major



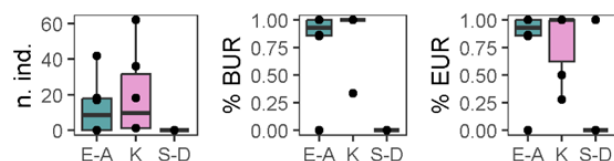
Trace



Microbial



Crustacean



274

275

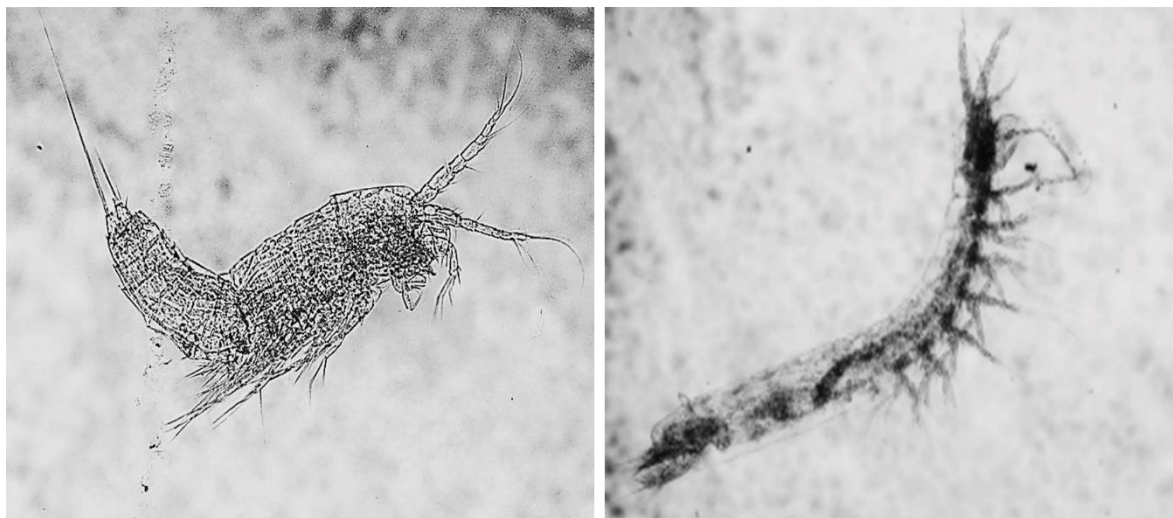
276 **Figure 2: Boxplots showing significant differences ($p < 0.05$, permutational t-test) in environmental and biological parameters**
 277 **among the three groundwater bodies (earth-alkaline in green; K-rich in pink, and sulfate-depleted in purple) of the Sabatini**
 278 **Mounts aquifer: Boxplot are arranged into five rows: field parameters, major chemicals, trace elements, microbes, and**
 279 **crustaceans. Each boxplot includes median lines, 25th-75th percentile boundaries, whiskers for 10th-90th percentiles, and black**
 280 **dots indicating outliers. BUR: burrowers; EUR: eurythermal.**



281 This microbial pattern was consistent over the two years (Table S1). Significant PERMANOVA outcomes were obtained for
282 microbial catabolic profiles such as those related to carbohydrates, polymers and aminoacids degradation among the three
283 GWBs, while no differences across the GWBs were detected for carboxylic acids and amines utilization. However, the
284 outcomes of polymers and aminoacids were biased by variance heterogeneity (Table S1), while those of carbohydrates were
285 not consistent over the two years of investigation, suggesting that the microbial functioning of the three GWBs did not
286 consistently differ in the catabolic profile based on carbohydrates were not consistent over the two years of investigation,
287 suggesting that the microbial functioning of the three GWBs did not consistently differ in the catabolic profile based on
288 carbohydrates. Overall, the catabolic profile of the microbial assembly of the three GWBs seem to be based on
289 carbohydrates, polymers and carboxylic acids, with variation over time, while the profiles based on amines and aminoacids
290 were much less represented (Table 1).

291 3.2 Crustacean assemblages

292 Out of the 20 biological samples analyzed, 8 (40%) did not contain any invertebrate specimens (Table A2e). We identified
293 210 crustacean specimens, belonging to 9 species (Table A2e). Seven of these taxa were groundwater-obligate dwellers
294 (94% of the total abundances; Table A2e) showing typical morphological adaptations to dark environments (Fig. 3).



295

296

297 **Figure 3: Groundwater-obligate crustacean species. The figure showcases two groundwater-obligate crustacean species found**
298 **within the target aquifer (Mounts Sabatini): the harpacticoid *Parapseudoleptomesochra italica* (0.73 mm in length, on the left) and**
299 **the syncarid *Meridiobathynella* sp. (0.8 mm in length, on the right). The specimens exhibit the typical morphological traits of**
300 **groundwater fauna: blindness, depigmentation, and hypertrophy of sensory appendages.**

301 Approximately 60% of the samples contained a single species only (Tables 1 and A2e). The class Crustacea Copepoda
302 dominated with 7 species, including 5 groundwater-obligate species, 6 species belonging to the order Harpacticoida, and 1
303 species to the order Cyclopoida. Three non-parametric estimators out of seven indicated that 100% of the expected



304 invertebrate biodiversity was collected during the sampling survey. However, the remaining estimators suggested that we
 305 likely missed out a minimum of 9% (Chao2) to a maximum of 21% (Jackknife1) of the expected biodiversity (i.e., from 1 to 3
 306 more species; Fig. S1).

307 PERMANOVA analysis indicated significant differences in the composition of crustacean assemblages; however, the
 308 outcome was affected by variance heterogeneity. On the other hand, the analyses revealed significant unbiased differences in
 309 the abundances of groundwater-obligate species among the three GWBs (Table S1), with the sulfate-depleted GWB lacking
 310 groundwater-obligate species (Tables 2 and S1; Fig. 2). Notably, the K-rich GWB lacked epigeal species. Remarkably, there
 311 were no significant differences observed between the two years (Table S1) suggesting the existence of stable and GWB-
 312 dependent assemblages over time. Overall, the K-rich and earth-alkaline GWBs exhibited comparable abundances and
 313 species richness (Table 2).

314

315 **Table 2: Taxonomic and functional composition of the crustacean assemblages in the three groundwater bodies (GWBs).**
 316 **Abundances (n. of ind) and other values are reported as mean values. Superscript letters (a, b, c) indicate statistical differences**
 317 **among GWBs (permutational post-hoc t-tests; $p < 0.05$). + indicates groundwater-obligate species. S-D: sulfate-depleted; E-A:**
 318 **earth-alkaline.**

319

Species abundance (n. ind.)	S-D	E-A	K-rich
* <i>Parapseudoleptomesochra italica</i> Pesce & Petkovski, 1980	0	7	17
* <i>Parastenocaris</i> sp.	0	1	0
* <i>Nitocrella psammophila</i> Chappuis, 1954	0	0	1
* <i>Pseudectinosoma reductum</i> Galassi & De Laurentiis, 1997	0	0	2
* <i>Acanthocyclops agamus</i> Kiefer, 1938	0	1	2
* <i>Meridiobathynella</i> sp.	0	5	0
* <i>Niphargus</i> sp.	0	0	2
<i>Elaphoidella gracilis</i> (Sars, G.O., 1863)	0	2	0
<i>Moraria poppei</i> (Mrázek, 1893)	1	0	0
Groundwater-obligate species	0 ^b	15 ^a	20 ^a
Trait locomotion			
Burrowers	0.0 ^b	0.8 ^a	0.9 ^a
Interstitials	0.1	0.0	0.0
Swimmers	0.0	0.0	0.1
Trait diet			
Fine sediments + microorganisms	0.1	0.5	1.0
Living microphytes	0.0	0.3	0.0
Trait feeding habit			
Deposit feeders	0.1 ^a	0.5 ^{a,b}	0.9 ^b
Collectors	0.0	0.0	0.1
Grazers	0.0	0.3	0.0
Thermal tolerance			
Eurythermal	0.1 ^a	0.8 ^{a,b}	0.8 ^b
Moderately stenothermal	0.0	0.0	0.1
Stenothermal	0.0	0.0	0.1

320



321 However, the taxonomic composition in the two GWBs was different. In detail, the groundwater-obligate harpacticoid
322 *Parapseudoleptomesochra italica* was the most abundant species, accounting for 67% of crustacean biodiversity in the
323 aquifer (Table A2e). The groundwater-obligate *P. italica* and *Acanthocyclops agamus* were found in both earth-alkaline and
324 K-rich GWBs (Table 2). *Nitocrella psammophila*, *Pseudectinosoma reductum* and *Niphargus* sp. (groundwater-obligate
325 species) occurred in the K-rich GWB, while *Parastenocaris* sp. and *Meridiobathynella* sp. (groundwater-obligate species)
326 were collected in the earth-alkaline GWB where the epigean *Elaphoidella gracilis* (epigean species) was also present (Table
327 A2e). The epigean species *Moraria poppei* was the only species found in the sulfate-depleted GWB, with a single individual
328 of *Moraria poppei* collected from well QA21 in 2014 (Table A2e).

329 The majority of the species collected in this study were burrowers and eurythermal. The locomotion and thermal tolerance
330 traits exhibited variation among the three GWBs, primarily driven by the higher percentages of eurythermal species in the
331 earth-alkaline GWB compared to the other two GWBs (Tables S1 and A2f; Fig. 2). Notably, moderately stenothermal and
332 stenothermal species were collected only from the K-rich GWB and never from the other two GWBs (Table A2f).
333 PERMANOVA outcomes indicated significant differences in the diet trait among the GWBs, although this result was biased
334 by variance heterogeneity (Table S1). The deposit feeding trait was the most prevalent among various feeding habits, with
335 collectors and grazers being less common, but the analyses did not identify any significant differences across the GWBs
336 (Tables 2 and S1). The pattern was consistent over time for all functional traits (Tables 2 and S1).

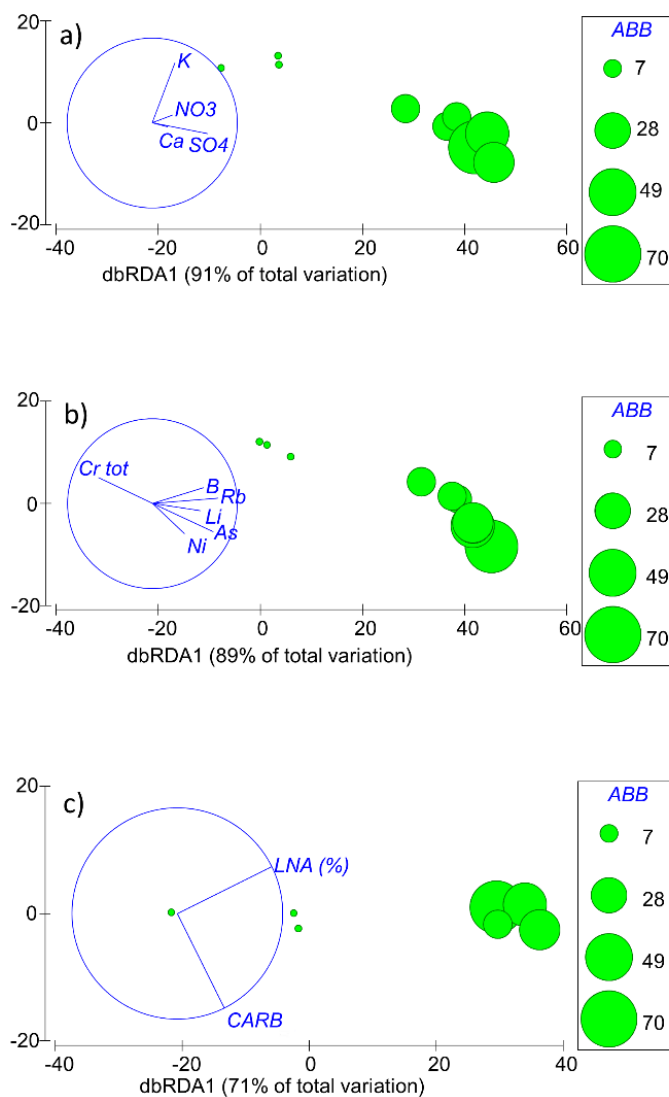
337 **3.3 Relationships between chemical and biological factors**

338 Concerning field factors, the multivariate models accounted < 65% of the variance of the taxonomic and functional
339 composition of the crustacean assemblages in the aquifer, with electrical conductivity, pH and ORP being the main
340 descriptors. None of the marginal models explained > 65% of the variance (Table S2).

341 Multivariate models based on major ions explained < 65% of the variance of the taxonomic composition, locomotion, diet,
342 feeding habits, and thermal tolerance traits (Table S2). However, four major ions (SO₄, Ca, NO₃, and K) together accounted
343 for 91% of the variance in groundwater-obligate species (Table S2), with abundances significantly decreasing in samples
344 depleted of SO₄ (Fig. 4). None of major chemical elements individually explained more than 65% of the variance (Table
345 S2). Concerning trace elements, multivariate linear models explained < 65% (Table S2) of the variance of the trait
346 locomotion and thermal tolerance. On the contrary, they explained > 65% of the variance of the taxonomic composition (best
347 model 71%: V, Cr, As, Se, and U), abundances of groundwater-obligate species (best model 89%: B, Al, V, Se, and Ba), trait
348 diet (best model 81%: Li, B, Cr tot, Ni, As, and U) and feeding habits (best model 71%: Li, B, Cr tot, Ni, As, and U), with
349 abundances and trait percentages increasing with increasing concentrations of all trace elements except for Cr tot (Fig. 4).
350 The contribution of individual trace elements to marginal models was always < 35%, when significant (Table S2). Finally,
351 concerning the microbial factors (TCC, LNA, HNA cells and catabolic pathways), conditional and marginal linear models
352 explained < 65% of the taxonomic and functional composition of the crustacean assemblages. However, LNA cells and the
353 carbohydrate pathways together explained 71% of the abundances of groundwater-obligate species (Table S2), with



354 abundances increasing with increasing LNA cells and microbes using carbohydrates as substrates (Fig. 4). The individual
355 contribution of LNA cells to the explained variance was 66%.
356



382 **Figure 4:** dbRDA plots showing the linear relations between abundances (ABB) of groundwater-obligate species and the factors
383 selected by the BEST procedure of the DistLM. a) major ions; (b) trace element and c) microbial factors (LNA%: percentages of
384 Low Nucleic Acid cells; CARB: carbohydrate catabolic profile).



385 **4 Discussion**

386 Despite recent advancements in the field, numerous knowledge gaps persist in identifying the factors that shape the
387 taxonomic and functional composition of groundwater crustaceans (Mammola et al., 2021). Particularly overlooked and
388 unexplored is the analysis of groundwater assemblages within volcanic GWBs and the role of trophic factors in shaping the
389 distribution of groundwater-obligate fauna through processes of environmental filtering (Saccò et al., 2019). To confront
390 these challenges, our study adopted a comprehensive, multidisciplinary approach that harnessed the collective knowledge
391 spanning hydrogeology, geology, microbiology and ecology.

392 In line with the generally stable nature of groundwater environmental conditions over time (Culver and Pipan, 2019), our
393 analyses confirmed and supported the occurrence of three adjacent GWBs, each characterized by distinct hydrochemical
394 facies and microbial community patterns. The sulfate-depleted GWB showed the highest mean temperature, the lowest
395 concentrations of SO₄, HCO₃ and U, the lowest microbial abundances and percentages of LNA cells, and the highest
396 percentages of HNA cells. The earth-alkaline GWB showed the lowest pH values and the highest electrical conductivity and
397 HCO₃ and Si concentrations, while the microbial abundances were comparable to those in the K-rich GWB. Finally, the K-
398 rich GWB presented the highest concentrations of Na, K, Li, B and Rb. We examined the crustacean assemblages of the
399 aquifer, revealing significant variations in both taxonomic and functional composition across the three GWBs. In detail, the
400 sulfate-depleted GWB lacked groundwater-obligate species, burrowers, and stenothermal or moderately stenothermal
401 species, while the K-rich and earth-alkaline GWBs, which showed comparable abundances and species richness, were
402 characterized by different species, which, however, showed the same functions concerning locomotion, diet and feeding
403 habit. Notably, stenothermal or moderately stenothermal crustacean species occurred in the K-rich GWB only, which was,
404 however, depleted of epigean species. We used a stringent cut-off criterion (Korbel and Hose; 2011, 2017; Di Lorenzo et al.,
405 2020) to identify the predictors of these differences, and concluded that the taxonomic composition is mainly driven by trace
406 elements such as V, Cr, As, Se, U. The main descriptors of the abundances of groundwater-obligate species are the major
407 ions SO₄, Ca, NO₃ and K, the trace elements B, Al, V, Se, and Ba, the microbial factors related to LNA cells and
408 carbohydrate catabolic profile. Finally, the trace elements Li, B, Cr tot, Ni, As, and U were the main drivers of the traits diet
409 and feeding habits. Water chemistry did not appear to exert a detrimental effect on the composition and functionality of
410 crustacean assemblages in the target aquifer, consistent with previous studies (e.g., Di Lorenzo et al., 2020), except for Cr.
411 This result can be attributed to the likelihood that the chemicals recognized as detrimental to groundwater organisms were
412 either not detectable or existed in concentrations that were not harmful in our study. For instance, trace elements, such as Ni,
413 Zn, As, Li, which are toxic to groundwater fauna at concentrations > 150 µg/L (Di Lorenzo et al., 2023), never observed in
414 the study area. Ammonium (which exhibited concentrations below instrumental detection limit in our study) is lethal to
415 groundwater-obligate species at concentrations > 12 mg/L, albeit causing cellular and physiological damage at
416 concentrations ≥ 36 µg/L (Di Lorenzo et al., 2015). Nitrate causes no harm to groundwater fauna at concentrations < 100
417 mg/L (Di Lorenzo et al., 2023), which are much higher than those observed in our study. Previous research has shown that



418 groundwater species are sensitive to pharmaceutical compounds, pesticides, and BPA (Di Lorenzo et al., 2023). In our
419 preliminary study (Ademollo et al., 2012), chlorinated pesticides were not detected and only traces of PAHs and PCBs were
420 found slightly above the detection limits (0.01 ng/L). High tolerance to SO₄ (which was positively correlated to groundwater
421 species abundances in this study) is known for many groundwater-obligate species of marine origin, such as *P. italica*, *N.*
422 *psammophila* and *P. reductum* (Galassi, 2001). For instance, *N. psammophila* has been collected from the chemoautotrophic
423 groundwater of the Frasassi cave system (Italy) where sulfate concentrations reach up to 199 mg/L (Galassi et al., 2017).
424 *Parapseudoleptomesochra italica* has been recorded from the Movable Cave (Romania) where groundwater has high sulfate
425 concentration and, similarly, *P. reductum*, has been discovered in the in sulfidic groundwaters of Melissotrypa Cave
426 (Greece) (Brad et al., 2021). While prior studies have reported synergistic toxic effects of pollutant mixtures on groundwater
427 species (Di Marzio et al., 2018), our research found no significant differences in chemical parameters among the various
428 GWBs that could account for varying toxicity levels. For instance, in the sulfate-depleted GWB, which lacked groundwater-
429 obligate species, most of the tested chemical elements exhibited lower concentrations compared to the earth-alkaline or K-
430 rich GWBs. Studies on the ecotoxicology of groundwater organisms have been limited by life history traits that make them
431 unsuitable for laboratory experiments (Di Lorenzo et al., 2019). Our study offered valuable insights into the sensitivity of
432 groundwater crustaceans to chemicals that have hitherto remained untested in prior research, including elements like
433 uranium and boron. The negative correlation between the abundances of groundwater-obligate species and Cr is, in our view,
434 not directly related to the toxicity of this element. Total chromium is present in the aquifer at concentrations ranging from
435 0.4 to 1.8 µg/L, which are considered harmless based on available literature data (Di Lorenzo et al., 2023). This might be a
436 statistical artifact since lower abundances of groundwater-obligate species are found in the sulfate-depleted GWB, where the
437 average Cr concentration is slightly higher than in the other GWBs. This is why we observe this negative correlation in
438 Figure 4. However, we believe that the absence of groundwater-obligate species in this GWB is related to microbial factors,
439 as we will explain below.

440 While the sulfate-depleted GWB lacks groundwater-obligate species, the earth-alkaline and K-rich GWBs differ in terms of
441 species composition. This outcome is consistent with previous studies. Iannella et al. (2020) observed that European
442 groundwater-obligate harpacticoid species (which represent 67% of the species in our study) were unable to disperse across
443 boundaries between two adjacent GWBs. Accordingly, Vaccarelli et al. (2023) observed that the dispersal of groundwater-
444 obligate copepod species in the Eastern Lessinian Massif (Italy) is constrained by non-fractured igneous rocks, as it appears
445 to be in our study. Only two groundwater-obligate species, *P. italica* and *A. agamus*, were found in both the K-rich and
446 earth-alkaline GWBs. The remaining species collected in this study were unique to one GWB only, such as the groundwater-
447 obligate *P. reductum* (a Tertiary relict of ancient marine origin; Galassi et al., 1999), which was collected from the K-rich
448 GWB, along with *N. psammophila*. The genus *Nitocrella*, which has direct marine origin, serves as an indicator of ancient
449 evolutionary events (Galassi et al., 2009). Notably, the K-rich GWB showed the highest concentrations of Li, B, and Rb,
450 which are characteristic elements of the deep geothermal facies and possibly of fossil marine waters of Neogene age (Duchi
451 et al., 2003). The presence of stenothermal and moderately stenothermal species in this GWB, and only in this one, suggests



452 that the habitat of the K-rich GWB is conservative and, therefore, suitable for preserving ancient evolutionary lineages
453 whose descendants survived in relatively stable habitats, and with no close relatives in surface environments, such as *P.*
454 *reductum*.

455 Six out of the nine indicators revealed that a small percentage of taxonomic diversity (ranging from 1 to a maximum of 2
456 species) was not captured in this study. This finding is commonly encountered because groundwater sampling is essentially a
457 blind process (Mammola et al., 2021). Wells serve as windows through which we gain insight into the subterranean
458 biodiversity in the portions of the aquifer surrounding them, but groundwater bodies extend more extensively (Ficetola et al.,
459 2019). However, we posit that the lack of groundwater-obligate species in the sulfate-depleted GWB is likely attributed to
460 the low percentages of LNA cells found in this GWB rather than to incomplete sampling. The LNA and HNA cell counts
461 exhibited variations among the three GWBs. Specifically, the sulfate-depleted GWB displayed the lowest LNA cell
462 percentages. As detected by flow cytometry, LNA and HNA cell groups are considered constitutive traits of aquatic
463 microbial communities, typically comprising cells of varying sizes, genome content, and phylogenetic affiliations (Gasol et
464 al., 1999; Proctor et al., 2018). Notably, HNA cells were recognized as an active fraction of the bacterioplankton community,
465 and their abundance was reported to positively correlate with heterotrophic production rates in freshwaters of different origin
466 (Rubbens et al., 2019). In contrast, LNA cells were traditionally thought to represent a more dormant or quiescent portion of
467 the aquatic microbial community (Lebaron et al., 2002). They were reported as small-sized microorganisms with slower
468 metabolic activity and a wide range of survival strategies suitable for thriving in adverse conditions, including oligotrophy
469 (Hu et al., 2022). In previous groundwater studies, LNA cells contribution and carbohydrate utilization were associated
470 under pristine and unbalanced nutrient conditions (Melita et al. 2019). This is explained by the role played by carbohydrates
471 as important energy-rich carbon source and storage molecules for the aquatic bacterial metabolism (Arnosti et al., 2014).
472 These observations highlight how groundwater quality can directly affect the functional properties of the aquatic microbial
473 communities with implications on the pattern of the energy fluxes among organic matter, microbes and the organisms
474 located in the upper levels of the food web. In this study, the high linear correlation ($r = 0.80$) between groundwater
475 crustacean abundance and LNA cells (supported by an explained variance $> 65\%$) suggested that groundwater crustaceans
476 could derive significant benefits from the presence of LNA cells. Considering the high abundances of deposit-feeders and
477 crustacean feeding on sediments and microorganisms in the earth-alkaline and K-rich GWBs, compared to the sulphate-
478 depleted GWB, we venture to speculate that they may selectively feed on LNA cells. If this hypothesis is confirmed, our
479 findings would offer supporting evidence that the feeding behavior of groundwater crustaceans can play an underrated role
480 in structuring the groundwater microbial community and biomass, with direct consequences on subterranean carbon turnover
481 and nutrient cycling.

482 Total cell counts also varied across the GWBs with the sulfate-depleted GWB showing the lowest values. However, we did
483 not observe a strong linear correlation ($r = 0.5$, $R^2 = 25\%$; Table S2) between TCC and groundwater crustacean abundance.
484 This result might seem counterintuitive since most of the collected species are known to feed on fine sediments and
485 microorganisms. Nevertheless, there should be a logical explanation for this outcome. The planktonic microbial community



486 represents a fraction of total aquifer microorganisms that is found in the interstitial water volume by detaching from
487 sediments (Flemming and Wuertz, 2019). Since the crustacean species examined in this study are not filtrators, it is probable
488 that they consume a relatively smaller portion of microbial planktonic cells when compared to sediment-attached ones. This
489 feeding habit could be the underlying cause of the weak correlation observed in this study. This observation raises the
490 possibility that a more robust correlation between diet and feeding habits and LNA cells might have emerged had we
491 included sediment-attached cells in our analysis.

492 On this matter, we did not observe a significant correlation between DOC and groundwater crustacean assemblages. This
493 suggests that DOC is likely not directly utilized by the species collected in this study but needs to be processed by bacteria
494 before becoming accessible to metazoans (Foulquier et al., 2009; Griebler and Lueders, 2009; Segawa et al., 2015).
495 Furthermore, DOC concentrations did not significantly differ across the three GWBs, being always <1 mg/L, in line with the
496 values normally observed in groundwater systems (Foulquier et al., 2010). DOC entering subterranean environments from
497 the surface primarily comprises stable and recalcitrant components that resist bacterial degradation, leaving only a small
498 fraction available for microbial communities (Shen et al., 2015). Our analyses indicated that the sulfate-depleted GWB had
499 the highest proportions of microbes metabolizing polymers, which are more complex and resistant to degradation than
500 simple carbohydrates or carboxylic acids (Oest et al., 2018; Melita et al., 2019). Additionally, we observed variations in the
501 proportions of microbes utilizing different substrates from year to year, likely influenced by surface-produced organic matter
502 types (Saccò et al., 2019; Melita et al., 2023). Our finding suggests that the sulfate-depleted GWB may lack the necessary
503 energy resources to sustain a resident crustacean assemblage. To provide a rough estimate, considering that a prokaryotic
504 cell contains approximately 25 fg of C (Griebler et al., 2002), it can be suggested that the average microbial biomass of $2.65 \cdot 10^4$
505 mg C/L recorded in the sulfate-depleted GWB might not be sufficient to sustain a community of groundwater-obligate
506 species.

507 **5 Conclusions**

508 Our multidisciplinary study delved into the taxonomic and functional composition of groundwater crustaceans and shed light
509 on the intricate dynamics of groundwater ecosystems. We unveiled significant variations in crustacean distribution and
510 functional traits across different groundwater bodies, with the sulfate-depleted groundwater body standing out as a unique
511 and seemingly inhospitable environment for groundwater-obligate species. Our findings pointed to the crucial role of
512 microbial communities in meeting the energy needs of groundwater-obligate crustaceans, thus indicating their potential
513 contribution to ecosystem services through a selective feeding strategy. Additionally, we provided insights into the
514 sensitivity of groundwater-obligate species to aquifer settings, expanding our knowledge of their tolerance to
515 hydrogeochemical and microbial factors. Our research underscores the importance of singling out diverse hydrogeological
516 contexts within individual aquifers. Potential avenues for future research encompass metagenomic studies on specific



517 microbial taxa that are at the base of groundwater food webs, while stable isotope analyses would help elucidate the dietary
518 preferences and food web dynamics of groundwater-obligate crustaceans and their impact on nutrient cycling.

519

520 **Data availability**

521 The raw data have been reported in Appendix A.

522 **Supplement**

523 The supplement containing Tales S1 and S2 related to this article is available online at:

524 **Author contribution**

525 TDL, SA, EP: Conceptualization; TDL, SA, MM, AZ, DP, SG, DR, ATDC, EP: Methodology; MM, AZ, DP, SG, DR,
526 DMPG: Data curation; TDL, SA, EP, DMPG: Writing- Original draft preparation. SA, EP, DMPG: Validation; TDL, SA,
527 EP, DMPG, ATDC: Writing- Reviewing and Editing.

528

529 **Competing interests**

530 The authors declare that they have no conflict of interest.

531

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728 **Appendix A**

729 **Table A1. Characteristics of the sampling sites within the three groundwater bodies (GWB).**

730

ID	Lat	Long	Elevation (m a.s.l.)	Depth (m)	Use	GWB
731 QA07	42.06812	12.50096	132	110	Domestic	sulfate-depleted
QA08	42.07234	12.51895	68	50	Domestic	earth-alkaline
732 QA09	42.07142	12.52757	49	29	Irrigation	earth-alkaline
QA13	42.09246	12.50991	187	115	Domestic	K-rich
733 QA15	42.08583	12.2710	146	115	Domestic	K-rich
QA21	42.08511	12.49676	176	105	Domestic	sulfate-depleted
734 QA24	42.05984	12.49368	107	80	Domestic	sulfate-depleted
QA25	42.08269	12.52501	126	120	Domestic	K-rich
735 QA29	42.09349	12.50305	145	64	Domestic	earth-alkaline
736 QA30	42.07626	12.50011	153	95	Domestic	sulfate-depleted

737

738 **Table A2. Physical-chemical, microbial, and crustacean data (taxonomic and functional) of the groundwater samples**
 739 **in the three groundwater bodies (GWBs) of Sabatini Mounts aquifer.**

740

741 **a) Field data (WT: depth to water table in m a.s.l.; ORP: oxidation-riduction potential in mV; T: temperature in °C;**
 742 **DO: dissolved oxygen in mg/L; EC: electrical conductivity in µS/cm).**

743

ID	YEAR	GWB	WT	ORP	T	pH	DO	DO%	EC
744 QA07	2014	S-D	65.0	211	18.0	7.6	7.1	75	490
745 QA08	2014	E-A	41.3	206	16.9	6.8	3.8	39	1048
QA09	2014	E-A	40.6	105	16.7	7.2	6.6	69	749
746 QA13	2014	K	112.5	194	16.2	7.5	8.5	88	675
QA15	2014	K	68.8	208	17.0	7.7	7.5	79	641
747 QA21	2014	S-D	104.1	137	18.6	7.7	5.0	54	479
QA24	2014	S-D	67.4	248	17.3	7.5	7.0	73	545
748 QA25	2014	K	64.4	244	17.1	7.8	5.0	52	640
QA29	2014	E-A	110.8	201	15.4	7.3	8.0	82	707
749 QA30	2014	S-D	82.2	221	17.4	7.5	8.1	85	525
750 QA07	2015	S-D	66.5	170	18.6	7.4	11.3	121	497
QA08	2015	E-A	41.2	229	17.5	6.8	3.9	41	1052
751 QA09	2015	E-A	40.3	58	17.0	7.4	6.9	72	746
QA13	2015	K	112.9	161	16.7	7.6	8.4	88	679
752 QA15	2015	K	68.6	260	17.0	7.4	6.9	73	645
QA21	2015	S-D	104.8	130	19.5	7.5	3.2	35	479
753 QA24	2015	S-D	67.7	170	17.4	7.3	8.1	85	551
QA25	2015	K	63.8	200	19.1	7.7	6.5	71	665
754 QA29	2015	E-A	111.3	141	16.7	7.3	6.1	64	711
755 QA30	2015	S-D	82.9	180	17.8	7.5	8.5	91	530

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762 **b) Major components (in mg).**

763

ID	YEAR	GWB	F	Cl	NO3	SO4	HCO3	Na	Mg	K	Ca	Si	DOC	
764	QA07	2014	S-D	1.6	20.9	7.1	4.8	237.9	30.1	8.1	41.5	39.4	25.3	1.5
765	QA08	2014	E-A	1.1	41.6	12.7	131.5	440.4	44.1	19.1	37.8	132.0	26.8	1.3
766	QA09	2014	E-A	1.0	17.6	44.6	28.3	358.7	25.1	5.2	29.6	32.7	26.7	0.8
767	QA13	2014	K	1.3	36.9	16.9	11.7	312.3	38.8	13.5	53.6	52.5	26.1	1.1
767	QA15	2014	K	2.1	31.0	17.5	18.6	269.6	49.5	10.7	53.3	39.5	23.2	0.8
767	QA21	2014	S-D	1.6	22.7	7.3	7.8	214.7	24.5	6.7	36.6	43.3	25.5	1.2
768	QA24	2014	S-D	0.9	32.9	20.5	7.6	241.6	27.2	10.4	32.3	51.4	27.8	0.8
768	QA25	2014	K	3.2	34.6	18.0	35.3	256.2	70.1	8.6	43.6	31.4	18.1	0.7
769	QA29	2014	E-A	0.6	29.6	25.1	21.9	331.8	28.3	11.9	31.8	81.7	26.8	0.9
769	QA30	2014	S-D	1.4	36.5	7.1	4.8	225.7	27.2	9.8	37.7	43.3	26.2	0.7
770	QA07	2015	S-D	1.6	20.4	7.0	4.6	244.0	15.6	7.1	38.2	33.7	18.9	0.5
770	QA08	2015	E-A	1.1	40.1	13.6	127.7	442.9	40.6	17.1	32.7	113.0	19.8	0.8
771	QA09	2015	E-A	1.0	17.0	33.3	27.2	363.6	27.2	10.2	31.9	77.8	16.1	0.9
772	QA13	2015	K	1.1	35.1	16.6	11.4	306.2	35.5	12.3	46.7	46.1	19.3	0.6
772	QA15	2015	K	2.1	30.7	16.7	18.5	278.2	32.9	10.1	48.4	36.1	18.0	0.8
773	QA21	2015	S-D	1.7	22.6	6.9	8.3	223.3	22.7	5.9	32.6	38.5	19.7	0.7
773	QA24	2015	S-D	0.9	29.4	18.2	7.2	242.8	24.7	8.9	27.7	43.5	19.9	0.7
774	QA25	2015	K	3.1	31.6	17.3	39.0	257.4	63.5	7.6	38.8	28.5	13.5	0.8
774	QA29	2015	E-A	0.6	29.5	25.8	20.4	328.2	26.5	10.4	27.7	68.8	20.2	0.5
775	QA30	2015	S-D	1.4	39.5	7.9	4.7	218.4	24.9	8.7	33.2	37.5	18.3	0.5

776

777 **c) Trace elements (in µg).**

ID	YEAR	GWB	Li	B	Al	V	Cr	Mn	Fe	Ni	Cu	Zn	As	Se	Rb	Sr	Ba	Pb	U
QA07	2014	S-D	20.1	76.1	11	40.2	1.8	2.7	34.8	0.4	0.5	17.8	20.1	0.52	28	560	74.1	0.1	2.9
QA08	2014	E-A	18.1	101	7.8	35.1	0.7	0.5	21.1	1.9	1.5	14.1	29.6	1.5	40	1396	104	0.2	30.7
QA09	2014	E-A	11.1	65.8	3.2	38.6	1.1	0.4	9.2	0.3	0.2	8.4	32.6	0.2	31	127	9.4	0.2	2.5
QA13	2014	K	23.6	111	13	38.1	1.1	0.9	23.2	0.5	0.8	24.7	13.2	0.2	29	849	102	0.2	6.1
QA15	2014	K	45.9	116	14	23.1	1.4	0.3	15.1	0.3	0.6	21.8	26.7	0.4	61	416	25.4	0.2	13.6
QA21	2014	S-D	22.3	65.4	2.9	29.5	1.3	0.9	21.2	0.9	0.4	13.1	31.7	0.4	19	489	68.5	0.2	4.8
QA24	2014	S-D	14.1	67.5	3.4	31.1	1.2	0.3	8.9	0.3	0.4	84.2	13.3	0.5	24	599	88	0.2	3.2
QA25	2014	K	70.5	144	12	24.5	0.6	1.4	26.4	59.4	0.7	22.4	43.1	0.2	64	331	35.7	0.3	17.8
QA29	2014	E-A	16.2	115	11	25.5	1.5	0.2	10.8	2.1	0.6	28.1	13.3	0.6	26	831	79.1	0.2	34.3
QA30	2014	S-D	19.7	71	5.1	30.1	1.2	0.2	11.5	0.9	0.3	32.2	20.3	0.2	19	593	72.1	0.2	2.5
QA07	2015	S-D	19.1	62.5	9.9	35.5	1.3	1.8	18.8	0.1	0.2	5.5	17.5	0.1	27	493	62.8	0.1	2.7
QA08	2015	E-A	17.7	91	9.7	31.3	0.6	0.4	15.2	0.2	1.3	2.2	31.9	1.5	37	1233	90.9	0.1	29.6
QA09	2015	E-A	14.5	66.2	4.1	18.3	0.6	9.6	11.0	0.1	0.2	3.6	15.7	0.7	22	578	129	0.1	15.1
QA13	2015	K	22.1	97	31	34.4	1.5	0.6	24.6	0.1	0.3	7.4	13.1	0.1	26	767	96	0.2	5.5
QA15	2015	K	44.9	111	9.6	22.1	1.1	0.2	11.2	0.1	0.6	6.4	27.9	0.3	57	401	25.1	0.1	11.8
QA21	2015	S-D	22.1	60.3	31	26.6	1.6	0.7	14.8	0.1	0.3	3.3	32.9	0.2	18	454	65.7	0.2	4.7
QA24	2015	S-D	13.6	57.3	16	27.8	1.1	0.5	61	0.1	0.3	104	12.1	0.1	21	527	82.5	0.6	2.8
QA25	2015	K	69.4	123	7.1	20.6	0.4	0.1	5.9	0.1	0.2	5.3	36.2	0.1	59	360	45.5	0.1	17.6
QA29	2015	E-A	15.5	98	31	22	1.8	0.3	11.2	0.1	0.4	8.9	12.3	0.1	23	717	76.5	0.2	27.9
QA30	2015	S-D	18.2	58.8	5.9	27.1	0.9	0.1	8.5	0.1	0.1	38.4	16.8	0.1	17	520	64.7	0.1	2.1

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783 **d) Microbial community characteristics. Total cell count (TCC in cell/mL), Low Nucleic Acidic and High Nucleic**
 784 **Acids cells are expressed as percentages of total absorbance (CARB: carbohydrates, POL: polymers; CARB_A:**
 785 **carboxylic acids, AM: amino acids, AMIN: amines).**
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ID	YEAR	GWB	TCC	LNA	HNA	HNA/LNA	CARB	POL	CARB A	AM	AMIN
QA07	2014	S-D	9896	66	34	0.5	40	25	20	11	4
QA08	2014	E-A	13950	80	20	0.2	31	14	27	23	6
QA09	2014	E-A	53423	81	19	0.2	37	7	26	25	6
QA13	2014	K	17705	65	35	0.5	21	41	20	18	0
QA15	2014	K	9315	70	30	0.4	31	27	28	14	0
QA21	2014	S-D	9283	68	32	0.5	40	25	20	11	4
QA24	2014	S-D	5409	63	37	0.6	40	25	20	11	4
QA25	2014	K	94954	80	20	0.2	31	27	28	14	0
QA29	2014	E-A	15265	67	33	0.5	46	13	25	17	0
QA30	2014	S-D	6291	61	39	0.7	52	18	19	12	0
QA07	2015	S-D	20307	67	33	0.5	0	95	0	0	5
QA08	2015	E-A	17182	82	18	0.2	36	13	25	18	8
QA09	2015	E-A	37164	77	23	0.3	40	11	23	22	4
QA13	2015	K	13148	66	34	0.5	10	25	55	6	3
QA15	2015	K	12447	75	25	0.3	0	34	58	0	8
QA21	2015	S-D	13822	72	28	0.4	0	69	29	0	2
QA24	2015	S-D	11843	62	38	0.6	0	51	47	0	2
QA25	2015	K	79904	75	25	0.3	0	30	70	0	0
QA29	2015	E-A	61110	67	33	0.5	44	22	16	10	8
QA30	2015	S-D	8175	67	33	0.5	0	60	40	0	0

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801 **e) Taxonomic composition. Pit: *Parapseudoleptomesochra italica*; Psp: *Paratenocaris* sp. Nps: *Nitocrella psammophila*;**
 802 **Pre: *Pseudectinosoma reductum*; Egr: *Elaphoidella gracilis*; Aag: *Acanthocyclops agamus*; Mpo: *Moraria poppei*; Msp:**
 803 ***Meridiobathynella* sp.; Nsp: *Niphargus* sp.; SB: cumulative abundances of groundwater-obligate crustacean species.**
 804

ID	YEAR	GWB	Pit	Psp	Nps	Pre	Egr	Aag	Mpo	Msp	Nsp	SB
QA07	2014	S-D	0	0	0	0	0	0	0	0	0	0
QA08	2014	E-A	12	2	0	0	0	0	0	3	0	17
QA09	2014	E-A	18	0	0	0	0	0	0	0	0	18
QA13	2014	K	1	0	0	0	0	0	0	0	0	1
QA15	2014	K	0	0	0	0	0	0	0	0	1	1
QA21	2014	S-D	0	0	0	0	0	0	1	0	0	0
QA24	2014	S-D	0	0	0	0	0	0	0	0	0	0
QA25	2014	K	62	0	0	0	0	0	0	0	0	62
QA29	2014	E-A	0	0	0	0	9	0	0	0	0	0
QA30	2014	S-D	0	0	0	0	0	0	0	0	0	0
QA07	2015	S-D	0	0	0	0	0	0	0	0	0	0
QA08	2015	E-A	13	0	0	0	0	2	0	27	0	42
QA09	2015	E-A	0	0	0	0	0	0	0	0	0	0
QA13	2015	K	1	0	0	0	0	0	0	0	0	1
QA15	2015	K	0	0	5	1	0	12	0	0	0	18
QA21	2015	S-D	0	0	0	0	0	0	0	0	0	0
QA24	2015	S-D	0	0	0	0	0	0	0	0	0	0
QA25	2015	K	36	0	0	0	0	0	0	0	0	36
QA29	2015	E-A	0	0	0	0	4	0	0	0	0	0
QA30	2015	S-D	0	0	0	0	0	0	0	0	0	0

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806 **f) Functional composition (in percentage): BUR: burrowers, INT: interstitial; SWI: swimmers; FS-M: fine sediments**
 807 **+ microorganisms; LM: living microphytes; D-F: deposit-feeders; COL: collectors; GRA_ grazers; EUR:**
 808 **eurythermal; MST: moderately stenothermal; STE: stenothermal.**
 809

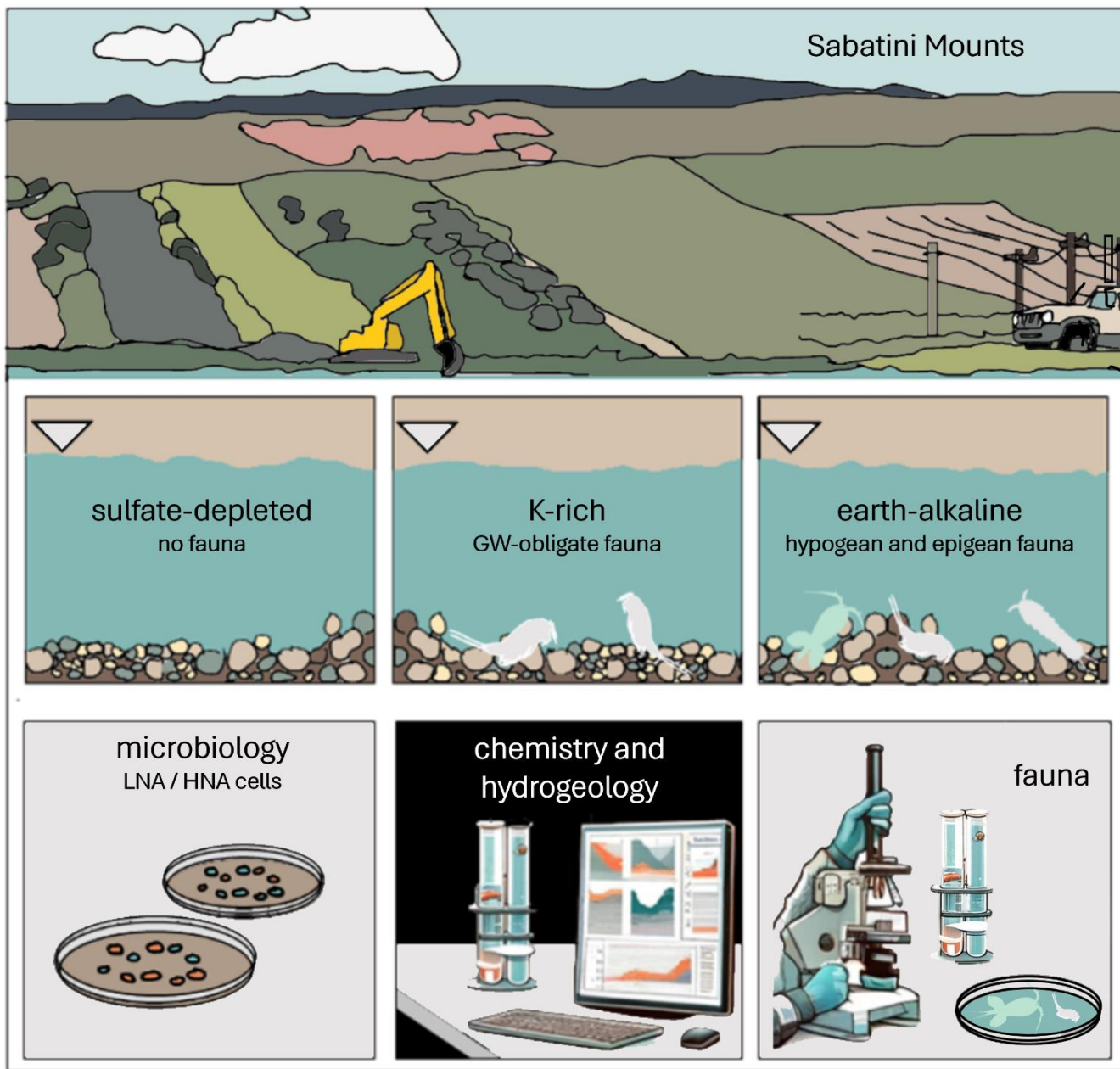
ID	YEAR	GWB	LOCOMOTION			DIET		FEEDING HABITS			THERMAL TOLERANCE		
			BUR	INT	SWI	FS-M	LM	D-F	COL	GRA	EUR	MST	STE
QA07	2014	S-D	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
QA08	2014	E-A	0.9	0.1	0.0	1.0	0.0	1.0	0.0	0.0	0.9	0.0	0.1
QA09	2014	E-A	1.0	0.0	0.0	1.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0
QA13	2014	K	1.0	0.0	0.0	1.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0
QA15	2014	K	1.0	0.0	0.0	1.0	0.0	0.5	0.5	0.0	0.5	0.0	0.5
QA21	2014	S-D	0.0	1.0	0.0	1.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0
QA24	2014	S-D	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
QA25	2014	K	1.0	0.0	0.0	1.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0
QA29	2014	E-A	1.0	0.0	0.0	0.0	1.0	0.0	0.0	1.0	1.0	0.0	0.0
QA30	2014	S-D	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
QA07	2015	S-D	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
QA08	2015	E-A	0.9	0.0	0.1	1.0	0.0	1.0	0.0	0.0	0.9	0.1	0.0
QA09	2015	E-A	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
QA13	2015	K	1.0	0.0	0.0	1.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0
QA15	2015	K	0.3	0.0	0.7	1.0	0.0	0.9	0.1	0.0	0.3	0.7	0.1
QA21	2015	S-D	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
QA24	2015	S-D	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
QA25	2015	K	1.0	0.0	0.0	1.0	0.0	1.0	0.0	0.0	1.0	0.0	0.0
QA29	2015	E-A	1.0	0.0	0.0	0.0	1.0	0.0	0.0	1.0	1.0	0.0	0.0
QA30	2015	S-D	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

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812 **Graphical abstract**



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