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Geological and hydrochemical prerequisites of unexpectedly high biodiversity in spring ecosystems at the landscape level

This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Cantonati M., Segadelli S., Spitale D., Gabrieli J., Gerecke R., Angeli N., et al. (2020). Geological and hydrochemical prerequisites of unexpectedly high biodiversity in spring ecosystems at the landscape level. SCIENCE OF THE TOTAL ENVIRONMENT, 740, 1-16 [10.1016/j.scitotenv.2020.140157].

Availability: This version is available at: https://hdl.handle.net/11585/904464 since: 2022-11-20

Published:

DOI: http://doi.org/10.1016/j.scitotenv.2020.140157

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Marco Cantonati, Stefano Segadelli, Daniel Spitale, Jacopo Gabrieli, Reinhard Gerecke, Nicola Angeli, Maria Teresa De Nardo, Kei Ogata, John D. Wehr, Geological and hydrochemical prerequisites of unexpectedly high biodiversity in spring ecosystems at the landscape level, Science of The Total Environment, Volume 740, 2020, 140157

The final published version is available online at: https://doi.org/10.1016/j.scitotenv.2020.140157

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Graphical Abstractive spring habitats as regional freshwater biodiversity hotspots



Hydrogeological characterization



Geodiversity







Extensive physical and chemical gradients

13 µS cm ⁻¹	Conductivity	11560 µS cm ⁻¹
1	Shading	5
0.2 mg L ⁻¹	Nitrates	48 mg L ⁻¹
0 µg L-1	Chromium	705 µg L-1

Impacts:

Water diversion, Intensive agriculture and industry in the plains.









High y biodiversity

- ► Study region: High geodiversity with diverse lithotypes and aquifer structures.
- ► Consequence: Wide variation in ecomorphological types and hydrochemistry.
- Effect: High landscape-level (γ) biodiversity.
- ► Conservation: Protection of representative and proportional regional groups of springs.
- ► Springs: Ideal systems in which to investigate geo-biodiversity relationships.

*Manuscript (double-spaced and continuously LINE and PAGE numbered)-for final publication Click here to view linked References

1 Geological and hydrochemical prerequisites of unexpectedly high

2 biodiversity in spring ecosystems at the landscape level

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19

20

22 ABSTRACT

This study explores the factors affecting the biodiversity of diatoms, vegetation with focus on 23 bryophytes, and invertebrates with focus on water mites, in a series of 16 spring-habitats. The 24 25 springs are located primarily from the mountainous part of the Emilia-Romagna Region (Northern 26 Apennines, Italy), and two pool-springs from agricultural and industrial lowland locations. Overall, data indicate that biological diversity (Shannon-Wiener, α -diversity) within individual springs was 27 28 relatively low, e.g.: *S*_{diatoms} = 0-46, *S*_{water-mites} = 0-11. However, when examined at the regional scale, 29 they hosted a very high total number of taxa (γ -diversity; $S_{diatoms} = 285$, $S_{water-mites} = 40$), including 30 several new or putatively-new species, and many Red-List taxa. This pattern suggested there is 31 high species turnover among springs, as well as high distinctiveness of individual spring systems. A 32 key goal was to assess the hydrogeological and hydrochemical conditions associated with this high 33 regional-pool species richness, and to provide a guide to future conservation strategies. There was a striking variety of geological conditions (geodiversity, captured mainly with lithotype and aquifer 34 structure) across the study region, which led to wide variation in the hydrosphere, especially in 35 36 conductivity and pH. Agriculture and industrial activities (anthroposphere) in the lowlands resulted 37 in nutrient enrichment and other forms of pollution. Across all 16 spring-systems, several 38 hydrogeological conditions most strongly influenced the presence or absence of particular biota and were determinants of species importance: spring-head morphology, hydroperiod, discharge, 39 current velocity, and elemental concentration. These findings have important practical 40 41 consequences for conservation strategies. Our data show that it is imperative to protect entire 42 regional groups of springs, including representatives of the different ecomorphological spring types, lithologies, and degrees of human influence. These findings suggest that springs, when 43 44 studied from an ecohydrogeological perspective, are excellent systems in which to further 45 investigate and understand geo-biodiversity relationships.

Keywords: Springs, Diatoms, Bryophytes, Zoobenthos, Water mites, Geodiversity

49 **1. Introduction**

50 Spring habitats possess remarkably distinctive ecological features and hold great importance for 51 biodiversity conservation; however, they are fragile ecosystems affected by many human impacts, 52 particularly water diversion and habitat destruction (e.g., Glazier, 2014). Springs in natural or near-natural 53 conditions have often been described as biodiversity hotspots (e.g., Cantonati et al., 2012a), but the type of 54 diversity and key factors have rarely been specified (but see e.g. Nascimbene et al., 2011; Pascual et al., 55 2020). Among the reasons often mentioned for spring habitat uniqueness and biodiversity are the marked 56 heterogeneity of water characteristics, their complex microhabitat mosaic structure, and their 57 characteristic ecotonal environment. Springs typically transition from groundwater to surface water, and 58 intergrade between aquatic and terrestrial habitats, leading to localized spatial heterogeneity (Cantonati et 59 al., 2012a). 60 The heterogeneous nature of spring habitats has been classified in a variety of ways (e.g., Glazier, 2014; 61 Cantonati et al., 2020). Seepages are known as the spring type richest in diatom and meiofauna species 62 (Cantonati et al., 2012a). Low-conductivity springs on siliceous (e.g., granitic) substratum, when not affected by acidification or other types of contamination, host very species-rich and distinct diatom 63 64 assemblages (e.g., Cantonati et al., 2009). Flowing springs are those most similar to streams, and indeed 65 often the transition from springhead to spring-fed streams is seamless, but nevertheless there are diatom 66 species with a marked preference for the uppermost section of the running-water system, but not in lower 67 reaches (e.g., Cantonati et al., 2012b). Limestone Precipitating Springs (LPS) or petrifying springs are one of 68 the few spring types indicated as priority habitat by the European Union Habitat Directive (EU-HD, 1992; 69 Cantonati et al., 2016), and shelter biota highly-adapted to this this special, limestone-precipitating 70 hydrochemical microhabitat. Biota in these systems include specific algae (in particular the bio-calcifying 71 desmid alga Oocardium stratum; Linhart and Schagerl, 2015). Inland-saline (mineral) springs are potentially 72 of high evolutionary significance as putative stepping-stone habits that facilitated the radiation in inland 73 waters of algal taxa of marine origin (e.g., Cantonati et al., 2019b).

74 Species composition and spatial distribution in springs is mainly determined by ion content, particlesizes of major substrata (determined by current velocity), shading, nutrients, and dissolved metals 75 76 (Bonettini and Cantonati, 1996; Cantonati et al., 2012b). Disturbances markedly alter spring-habitats and 77 can result in clearly-detectable changes in diatom assemblages, in particular by alterations to the physical 78 structure of springs and contamination with nitrates (e.g., Angeli et al., 2010). Diatom Red List species 79 (percentage of species in threat categories) were shown to react sensitively to habitat alteration and 80 destruction (e.g., water capturing), as well as to nitrate pollution (Cantonati et al., 2012a). The recent 81 (Hofmann et al., 2018) publication of an updated and expanded version of the diatom Red List for Central 82 Europe (first edition: Lange-Bertalot, 1996) allows for an even more effective use of this tool for spring-83 habitat integrity evaluation in this geographic region.

84 Spring-ecosystem hydrochemistry is influenced by a variety of pre- and post-emergence factors affecting 85 the aquifer, including geology (e.g., Cantonati et al., 2020), climate, vegetation, and land-use (Merk et al., 86 2020). The close relationship between spring-dwelling communities and geological formations of the 87 aquifers has been clearly demonstrated for diatom assemblages (Werum and Lange-Bertalot 2004). In 88 some cases, the occurrence of individual diatom species appears to be correlated with the dominant 89 lithotype in the drainage basin, such as dolomite or granite. The occurrence of some diatom species in 90 spring ecosystems has been studied in detail and their ecological characterization often shows that their 91 distribution is driven by a complex combination of hydrogeological and chemical factors. For instance, 92 Achnanthidium dolomiticum (Cantonati and Lange-Bertalot, 2006), as the specific epithet suggests, is found 93 on dolomite $[CaMg(CO_3)_2]$ substrata. However, Cantonati et al. (2012b) found that this species is 94 characteristic of karstic springs with strongly fluctuating discharge. Recently, A. dolomiticum was also found 95 in springs emerging on other rock types, but were nonetheless influenced by groundwater enriched in 96 magnesium (Cantonati et al., 2017).

Bryophytes in spring habitats, like diatoms and other photoautotrophs, are strongly influenced by the
geological characteristics of the substratum (e.g., Nascimbene et al., 2011). Among invertebrates, water
mites are apparently a group with among the greatest number of spring-dependent species; this points to a

particularly strong relationship between this group and conditions in the spring environment (e.g., Gereckeet al., 2018).

102 Despite these patterns in individual studies, few papers have addressed the broader relationship 103 between biota and spring hydrogeology (e.g., Van der Kamp, 1995; Cantonati et al., 2020). The link 104 between geological diversity, or "geodiversity" (e.g., Gray et al., 2013) and biodiversity is increasingly 105 gaining interest, but far more studies are needed to provide empirical evidence for this connection 106 (Alahuhta et al., 2020). The role of geodiversity in positively influencing biodiversity has been recognized 107 for terrestrial ecosystems (e.g., Bailey et al., 2017), but studies on freshwater habitats are still rare (e.g., 108 Kärnä et al., 2019, who worked on streams). The Northern Apennines (Italy) are renowned as a highly 109 complex geoscape and area of high geological diversity (e.g., Castellarin, 2001; Boccaletti et al., 2004; 110 Marroni et al., 2010). Therefore, we focused this study specifically in the Emilia-Romagna Region for a 111 deeper understanding of spring-habitat biodiversity and a better understanding of their ecological and 112 geological drivers. We examined the hydrogeological and hydrochemical conditions associated with high regional-113 pool species richness and landscape-level (y) biodiversity with the goal to show how high 114 115 lithological and aquifer-structure diversity generate a multiplicity of ecomorphological and

116 hydrochemical environmental settings, and how this translates into high y diversity. We focused

our studies primarily on two groups of organisms, diatoms and water mites, which have

118 contrasting ecological roles within the spring ecosystem. The consequences of our findings for

- spring-habitat conservation strategies are highlighted.
- 120
- 121
- 122 **2. Methods**

We examined sixteen springs: Names and codes (with explanations) are listed in Table 1, and will be used throughout the paper. Data from these springs are listed by increasing mineral content, from lowest to highest conductivity.

Springs were selected on the basis of the following criteria: (1) location in different types of nature reserves; (2) representativeness of the diverse ecomorphological and hydrochemical spring types, and of the main lithotypes occurring in the study area; (3) availability of medium-term data series; (4) location within the main northern Apennine aquifer types; (5) location in the different altitudinal belts; (6) presence of a permanent hydraulic regime; and (7) in natural or near-natural conditions. Seventy separate morphological, physical, and chemical variables (Supplementary Material Table 1) and variables were determined for each of 16 selected springs.

133

134 2.1. Sampling surveys

135 Field work was carried out in the summers of 2011 and 2012 (hydrochemistry and biota).

136 Hydrogeological observations and monitoring of selected springs were performed continuously within a 6-

137 year timeframe, from 2007-2013. The position of the springs was recorded by measuring the geographic

138 coordinates with a GPS (Garmin eTrex 10; Garmin Ltd., Olathe, KS, USA). Geomorphology (configuration)

139 was assessed recording the main characteristics on sampling forms and drawing sketches. Current-velocity

140 was assessed using the five-point scale reported in Cantonati et al. (2012b). Canopy cover (as a measure of

141 shading) was assessed with a hemispherical densiometer.

142 Depending on local conditions, discharge measurements were performed either using the volumetric

143 method (with low flow rates) or using flumes and weirs to convey all the water inside a graduated

144 container. Measures were repeated at least three times at each site. Temperature, specific conductance at

145 25 °C, and pH were measured on-site by means of handheld Eutech temperature-conductivity meter (Cole-

- Parmer, Vernon Hills, IL, USA) along with each discharge measurement. Finally, at the Poia_MiSp-SC and
- 147 Laba_LPS-le springs, discharge values were determined with an STS digital pressure transducer (model

148 DL/N-64, STS - Sensor Technik Sirnach, Sirnach, Switzerland), whereas temperature and conductivity were

measured with a digital thermometer-conductimeter (Hanna Instruments; Woonsocket, RI, USA). To
calculate variability indices, discharge, temperature, and electrical conductivity data collected in the field
during the EBERs project have been integrated with own (SS) datasets or available from the literature
(Chiesi and Forti, 2009; Filippini, 2009-2010; Gherardi, 2011-2012).

153

154 2.2. Geology and Hydrogeology

155 Geological data used for this work was derived from the database of the geological map of the Emilia-156 Romagna Region, scale 1:10,000 (Geological cartography of Appennines, scale 1:10,000, 2012).

157 A variable termed 'lithology' was defined to express the main geological subdivisions in the Northern

158 Apennines and their influence on the mineral content of the studied spring waters. Lithological classes

were ordered on the basis of the degree of solubility of the rock and therefore on the basis of their capacity

160 to condition the chemistry of the waters during flow inside the aquifer. 'Lithology' expressed the range of

161 conditions from poorly-soluble rocks (e.g. siliciclastic arenites and basalts) to karst rocks composed of

162 evaporatic Triassic - Messinian rocks, and classified from 1 to 8, as follows: (1) siliciclastic arenites, (2)

163 ophiolites (basalts), (3) ophiolites (peridotites), (4) ophiolites (serpentinized peridotites), (5) lithic arenites,

164 (6) limestone-marly flyschs and calcarenites, (7) alluvial deposits, and (8) Triassic and Messinian evaporites

165 (details in Table 1).

166 The springs were classified on the basis of variation in discharge through time, and quantified using the

167 variability index (Rv) introduced by Meinzer (1923). This index is a function of the maximum (QM),

168 minimum (Qm), and mean (Qmed) discharge values within the hydrological year: Rv = ((QM -

169 Qm)/Qmed)*100). Values of Rv<25% indicate springs with constant discharge, whereas Rv between 25%

and 100% are defined as sub-variable. Higher values (Rv > 100%) identify springs with variable discharge.

171 This index was also implemented for temperature and specific conductance data. For several springs

172 (Helo_LCSe, CaLS_LCFS, GeRi_MiSp-Su, FoVR_PS-Sh, and FoVR_PS-Su), the Rv values had to be estimated.

173

174 2.3. Hydrochemistry

175 Sampling was conducted using polyethylene (PE) bottles previously cleaned with ultra-pure HNO₃

176 (Ultrapure grade, Romil, Cambridge, UK), and then rinsed several times with ultra-pure water (Purelab Ultra

177 Analytic, Elga Lab Water, High Wycombe, UK) to minimize the risk of external contaminations. Water

samples for major ions and nutrients were kept chilled (ca. 4 °C) in fridges and fridge bags until analysis.

179 Detailed hydrochemical analyses were carried out following standard methodology (APHA, 2000). Ca²⁺,

180 Mg²⁺, Na⁺, K⁺, NH₄⁺, Cl⁻, NO₃⁻, SO₄²⁻, Br⁻, F⁻ were analysed by ion chromatography (ICS 1500 Dionex Corp.,

Sunnyvale, CA, USA), and dissolved nutrients $(N-NO_2^-, N-NH_4^+, P-PO_4^{3^-}, TP, TN, Si)$ by standard absorption spectrometry (details in Cantonati et al., 2012b).

183 The bottles of water samples for trace elements were capped after sampling, packed in double LDPE 184 bags, and transported to the IDPA-CNR Laboratories (Milan, Italy), where samples remained frozen until 185 they were analysed. The samples were thawed at room temperature under a class 100 laminar flow clean 186 bench, and 10 mL aliquots were transferred to 12 mL ultra-clean LDPE vials and acidified with ultra-pure 187 HNO₃ to obtain 2% solutions (v/v). Concentrations of 29 elements (Li, Be, Na, Mg, Al, Ti, V, Cr, Mn, Fe, Co, 188 Ni, Cu, Zn, Ga, As, Rb, Sr, Mo, Ag, Cd, Sn, Cs, Sb, Ba, Tl, Pb, Bi, U) were determined by Inductively Coupled 189 Plasma Sector Field Mass Spectrometry (ICP-SFMS; Element2, Thermo-Fisher, Bremen, Germany) equipped 190 with a desolvation system (APEX IR, Elemental Scientific, Omaha, US). The system was calibrated by 191 external calibration using a certified multi-standard solution containing the 29 trace elements (IMS-102, 192 UltraScientific, US). Concentrations in the standard solutions ranged from 0.005 to 50 ng/g for all the 193 elements except Na, Mg, Al, Fe, Mn and Ba which were calibrated at higher concentration values (from 1.0 194 to 1000 ng/g). The accuracy was evaluated (ION 96.2, National Water Research Institute, Canada) and 195 ranged from 76% (Al) to 105% (Sn). Average precision was always better than 12%.

In order to evaluate the relative contribution from natural (e.g., rock and soil dust) versus anthropogenic sources, trace element concentrations were expressed in the form of crustal enrichment factors (EF_{Ba}). EF_{Ba} is defined as the concentration ratio of a given element to that of Ba (or any other conservative element which derives mainly from rock and soil dust), normalized to the same concentration ratio characteristic of the upper continental crust (after Wedepohl, 1995). However, given the large variations in the composition of rock and soil, enrichment factors within \pm 10 times the mean crustal abundance (i.e. EF_{Ba} values ranging from ~ 0.1 to 10) do not likely demonstrate the input from sources other than rock and soil dust. Conversely, any EF_{Ba} value significantly greater than 10 strongly suggests contributions from other natural sources and/or anthropogenic sources. Elements with EF_{Ba} values between 10 and 100 are considered moderately enriched, indicating one or several other sources in addition to the crustal material. Finally, elements with EF_{Ba} values greater than 100 are considered highly enriched, suggesting a severe anthropogenic contribution (after Veysseyre et al., 2001).

208

209 2.4. Diatom sampling, identification, and quantification

210 Diatom assemblages were sampled and treated following the methods described previously for spring 211 habitats by Cantonati et al. (2012b), with specific designation of the spring-head area (= eucrenal, defined 212 here as the submerged portion at the time of sampling and the inner bank clearly influenced by the water 213 during the year), choice of substrata, and sample treatment. Epilithic diatoms were collected by brushing 214 ten stones. For diatoms epiphytic on the dominant bryophyte species, the epibryon were collected from 215 the most frequent and abundant bryophytes in each sample location (see Supplementary Material Table 4). 216 The collected materials, including the bryophytes, were cut into small pieces and were digested using 217 hydrogen peroxide (Cantonati et al., 2012b). The cleaned material was mounted in Naphrax (refractive 218 index of 1.74). For each sample, three cover-slips were prepared on one permanent slide, and a pooled 219 total of about 450 valves were counted. All samples (original samples, suspensions of prepared material, 220 and permanent mounts) have been catalogued and deposited in the collections of the MUSE – Museo delle 221 Scienze (Trento) (access codes: cLIM007 DIAT 1956-2002) along with information about the abundance of 222 the species found and the main environmental variables.

Counting was conducted with a Zeiss Axioskop 2 at 1000x magnification (Zeiss, Oberkochen, Germany).
The most updated taxonomy and nomenclature available at the time when counts were performed was
applied (in particular, Levkov, 2009; Hofmann et al., 2011; Lowe et al., 2014), and other identification
reference works followed were as in Cantonati et al. (2012b). Nomenclature and some taxonomic concepts

were updated using Cantonati et al. (2017), AlgaeBase (Guiry and Guiry, 2020), DiatomBase (Kociolek et al.,
2020), Diatoms of North America (Spaulding et al., 2019), the Freshwater Diatom Flora of Britain and
Ireland (Jüttner et al., 2020). To confirm identifications and document taxa with poorly-observed
ultrastructure, several taxa were examined with SEM (Zeiss-EVO40XVP, Carl Zeiss SMT Ltd., Cambridge, UK)
at the MUSE - Museo delle Scienze (Trento) or at the University of Frankfurt using a Hitachi S-4500 (Hitachi
Ltd., Tokyo, Japan) at high vacuum on gold-coated stubs.

233

234 2.5. Bryophytes and vascular plants

Within the eucrenal, all the bryophytes and vascular plants were identified and assigned a score according to their visually estimated percent cover: 1 = < 5%, 2 = 5-25%; 3= 25-50%; 4 = 50-75%; 5 = 75-100 (e.g., Braun-Blanquet, 1964; Tomaselli et al. 2011). Species nomenclature follows Aeschlimann et al. (2004) for vascular plants, Hill et al. (2006) for mosses, and Ros et al. (2007) for liverworts. Specimens of vascular plants were collected as needed to confirm field identifications at a later stage. Bryophytes were systematically collected and will be deposited in the Herbarium of the MUSE – Museo delle Scienze, Trento, ltaly (Code = TR).

242

243 2.6. Zoobenthos

244 Cumulative sediment samples from all significant substrata were preserved in 70% denaturated ethanol or in 4% formaldehyde; animals from the latter was later transferred to 70% denaturated ethanol. During 245 246 sorting, water mites were then preserved in Koenike's fluid (glycerol:acidic acid:distilled water 10:3:6). 247 Before sorting, samples were washed and divided into two fractions (>1.000 μ m and <1.000 μ m). The large-248 scale fraction was investigated at lower magnification, the fine fraction at high magnification under a Zeiss 249 SV6 stereo microscope. All samples were examined completely; individuals encountered were counted 250 from the whole sample, although not all were directly enumerated. In cases of high densities, a quarter of 251 each dish was counted and the resulting number multiplied by 4. Resulting specimen numbers were 252 rounded and were used for calculating categories of relative density for all taxonomic groups recognized.

253 The material was sorted at the taxonomic level of orders, or, in the case of Diptera, families. As a general 254 rule, for taxa in low densities, all specimens were sorted and enumerated, while for more frequent taxa, 255 only a representative number of specimens was preserved. During treatment, attention was paid to 256 obvious differences in morphology and size classes, trying to sort out different morphotaxa in relation to 257 their relative abundance. In order to get insight into the whole diversity spectrum (often reflected by 258 differences in body weight and consequently in sedimentation rate), specimens of all taxa were selected 259 during the whole sorting process, from the first to the last Petri dish. For each spring, and each 260 order/family, a tube with two labels (taxon name and site abbreviation; information on collecting site and 261 date) was deposited in the collection. A part of the material has been sent to specialists, the rest was 262 deposited in the collections of the MUSE– Museo delle Scienze (Trento) Limnology Section (access codes: 263 cLIM007).

264

265 2.7. Data processing and statistical analyses

For all diatom species collected in this study, a threat status (a measure of rarity) was assigned,
according to current (Hofmann et al., 2018) and previous Red List data (Lange-Bertalot, 1996). For the
species present in both lists, a check was made if the conservation status is improving or declining.
Hofmann et al. (2018) provide further ecological attributes (trophic and mineralization preferences, aerial
species) used in this study (Supplementary Material Table 3).

271 Shannon-Wiener diversity (Shannon, 1948) was calculated using a base-2 logarithm. Canonical 272 Correspondence Analysis (CCA) was calculated with the package vegan (Oksanen et al., 2019) in the R 273 statistical environment (R CORE TEAM, 2018). CCA was selected over other ordination techniques, such as 274 redundancy analysis (RDA), after evaluation of the length of the gradient. Model selection was performed 275 with the orddistep function of vegan, with an automatic, stepwise model with constrained ordination. Only 276 the environmental variables significant in the last step of selection was then used for the final CCA model. 277 Significance of the CCA model, terms, and constrained axes was tested with an ANOVA like permutation 278 test (999 permutations).

A Mantel test was used to assess correlation between epibryon and epilithon diatoms and vegetation. The significance of the statistic is evaluated by permuting rows and columns of the first dissimilarity matrix. The function to perform the Mantel test is available in the vegan package. To study α , β , and γ diversity for diatoms, we used an additive diversity partitioning, where the mean values of α diversity at lower levels of a sampling hierarchy are compared to the total diversity in the entire data set (γ diversity, Crist et al., 2003); this function is available in the vegan package.

- 285
- 286

287 **3. Results**

288 3.1. Ecomorphology, Geology and Hydrogeology

289 The ecomorphological classification of the springs studied here led to the recognition of seven spring-290 types, listed in order of increasing mineral content of the spring water (Table 1, Fig. 3). The spring codes are 291 explained in Table 1, and include, after the underscore, an abbreviation of the spring type, and, after the 292 hyphen, an abbreviation of the possible sub-category. Within shaded, flowing springs, a subgroup was 293 characterized by the occurrence of the macroscopic crustose red alga Hildenbrandia rivularis. 294 The geological (lithological) and hydrogeological classification of the studied springs is provided in Table 295 1. Our study also included two lowland pool springs in alluvial deposits, locally called *fontanili*. Hard Rock 296 Aquifers (HRA) is the best represented group because the ophiolites, and in particular the turbiditic units, 297 both silico-clastic, and calcareous and calcareous-marly, cover very vast and seamless territories (Fig. 1).

298 *3.2. Hydrochemistry*

The study revealed several factors that ranged widely among spring-types (Fig. 4, Supplementary Material Table 1, Table 2). The conductivity gradient of the studied springs (13-11560 μS cm⁻¹) spanned more than three orders of magnitude. In one system, Prin_ShFS-pH, very high pH (11.2) was due to enrichment with sodium and chloride. The two mineral springs (Poia_MiSp-SC, GeRi_MiSp-Su) were characterized by high sodium, chloride, sulphate, and total phosphorus (TP) values. TP enrichment is otherwise rare in most systems, even in the agricultural lowland limnocrenes (*fontanili*) contaminated by
nitrate and chloride (Table 2). The MtNe_ShFS-Hi (aquifer: deep-seated gravitational deformations in slopes
- DSGSD) and CiLi_HygS springs, both coming to daylight on ophiolithic rocks, were enriched with sodium,
magnesium, chromium, nickel, arsenic, uranium, molybdenum, cadmium, antimony, titanium
(Supplementary Material Table 2, Table 2).

309

310 *3.3. Diatoms*

311 In the spring GeRi MiSp-Su, bryophytes were not found (see below), and diatoms were absent also on 312 lithic material. Diatom analyses of samples from the remaining 15 springs revealed a total species richness 313 of 285 taxa (272 identified to the species -or intraspecific- level) belonging to 63 genera (Supplementary 314 Material Table 3). Red-List threat-category data were available for nearly all the taxa identified to species 315 level: only nine taxa were not listed (Hofmann et al., 2018). A high proportion (60%) of the species for 316 which Red List information was available were found to belong to one of the key threat categories (1, 2, 3, 317 G, R, V, D oligotraphentic) (Supplementary Material Table 3). A selection of frequent and abundant, 318 characteristic, rare, and Red-List species is summarized in Fig. 5. 319 Two of the species identified were listed in the highest threat category, i.e. threat category 1 320 ("threatened with extinction"): Achnanthidium trinode (Fig. 5 11b) and Eunotia cisalpina (Fig. 5 2b). Several 321 more species in the studied springs are members of threat category 2 ("strongly threatened"), including 322 Achnanthidium dolomiticum, A. rosenstockii, Brachysira calcicola, B. vitrea, Cymbella tridentina (Fig. 5 12a), 323 C. tumidula (Fig. 5 12b), Cymbopleura austriaca, Encyonema hebridicum, Encyonopsis lange-bertalotii, 324 Eucocconeis flexella, Eunotia kruegeri, E. nymanniana, E. palatina, E. sudetica, E. tetraodon, Kolbesia 325 carissima (Fig. 5 2d), Navicula angusta (Fig. 5 1a), N. dealpina, Psammothidium altaicum, Rossithidium 326 anastasiae, R. pusillum, Sellaphora stroemii.

On the basis of in-depth studies on LM morphology, plastid shape and arrangement, SEM ultrastructure,
 ecology and distribution, and extensive literature research, eight species in the genera *Achnanthidium*,

Amphora, Delicata, Eunotia, Fragilaria, Halamphora, Navicula, and Planothidium were recognized as new
 to science (Cantonati et al., 2019a;b; MC, Horst Lange-Bertalot, unpublished data).

331 Ecological preferences were assigned (based on Hofmann et al. 2018) for 252 of the 272 taxa identified 332 at least to the species level (Supplementary Material Table 3). Of these only 24 species (= 9.5%) were 333 classified as aerial, and included Cymbopleura austriaca, Delicata minuta, Diploneis minuta, Encyonema 334 alpinum, E. bipartitum (Fig. 5 9f), Halamphora montana, H. normanii, Hantzschia amphioxys, almost all 335 Humidophila spp. found (excepting H. schmassmannii), Hygropetra balfouriana, Nitzschia harderi, Nupela 336 lapidosa, Pinnularia obscura, Simonsenia delognei, Surirella terricola, Tetracyclus rupestris, Tryblionella 337 debilis. Most species were encountered in only one or two springs. Table 3 shows richness (S) and Shannon-338 Wiener (H') ranges. Species richness in the epilithon ranged from 7 (Poia_MiSp-SC) to 46 (CaLS_LCFS); 339 richness in the epibryon ranged from 5 (Poia_MiSp-SC) to 65 (Helo_LCSe). The variable 'lithology' was 340 correlated significantly with both Shannon-Wiener diversity of epibryon diatoms (r = -0.74; p = 0.003) and

341 vegetation (*r* = - 0.78; *p* < 0.001, Fig. 6).

A CCA (Table 4, Fig. 7) indicated that the distribution of the sites and characteristic species was mainly 342 343 determined by differences in chromium, nitrates, elevation (for epilithon), and by nitrates, magnesium, 344 bicarbonate alkalinity, and elevation (for epibryon). This is confirmed by the placement of typical mountain 345 species (Psammothidium daonense, Achnanthidium trinode (Fig. 5 11b), Nupela lapidosa, Gomphonema 346 tenoccultum (Fig. 5 11c), Odontidium neolongissimum) close to the apex of the elevation vector. The 347 eutraphentic Fragilaria famelica is positioned close to the head of the nitrate vector, and indeed this 348 species was abundant in FoVR_PS-Su, where the highest nitrate values were measured (Table 2). 349 The total species richness (y-diversity) recorded across all springs was partitioned into the average 350 number of species that occur within a sample (α) and the average number of species absent from a sample 351 (β), Table 5. The partition showed that α diversity was on average 23.5 in the epilithon and 28.4 on average 352 in the epibryon, each of which is relatively low, compared to total y-diversity (199 and 214 species).

353

354 *3.4. Bryophytes and vascular plants*

355 A total of 93 macrophytic plant species were observed across all sites (43 bryophytes, and 49 vascular 356 plants; Supplementary Material Tables 4-5), with an average richness of 9.6 per site. In the spring GeRi_MiSp-Su both bryophytes and vascular plants were absent, whereas the richest spring was Helo_LCSe, 357 358 with 13 and 15 species respectively for bryophytes and vascular plants. Of note were the bryophytes 359 Hygrohypnum eugyrium (the second record in Italy), Fissidens viridulus, Racomitrium aciculare, and 360 Southbya tophacea; taxa that were not recorded after 1950 in the Emilia-Romagna Region. Most species 361 were encountered only once or twice. Species observed more than 5 times are common in many aquatic 362 habitats, such as Bryum pseudotriquestrum, Palustriella commutata, and Brachythecium rivulare. Other less 363 frequent species occurred in particular spring types, including Didymodon tophaceus and Eucladium 364 verticillatum in LPS springs, and Sphagnum spp. in seepages. The vegetation composition, designated here 365 as plant and bryophyte assemblages together, was highly correlated with the epibryon diatom assemblages 366 (Mantel statistic r = 0.95; p < 0.001), and correlated, but less so, with the epilithic diatoms (Mantel statistic 367 r = 0.30; p < 0.011).

368

369 3.5. Zoobenthos

370 *3.5.1. Overview on invertebrates*

371 The invertebrate groups encountered and the numbers of specimens counted, are given in 372 Supplementary Material Table 6. The most abundant groups were chironomids, followed by ostracods, 373 mites, and stoneflies. The total specimen number in samples collected during the investigation exceeded 374 ten thousand. A considerable abundance and frequency of groundwater taxa (= stygophilous) were 375 observed. Among these, particularly common were amphipods in the genus Niphargus, stygobiont 376 hydrobiid snails, and the mites genera Stygothrombium and Frontipodopsis, the latter two taxa previously 377 not published from spring sites in Italy. An interesting finding was also a larva of the psephenid beetle 378 genus Eubria, a taxon with a very scattered distribution in Europe. The relationship between selected 379 groups of invertebrates and lithology is shown in Fig. 8. Of these, only chironomid species composition was 380 significantly correlated to lithology (r = -0.65; p = 0.006).

381

382 3.5.2. Water mites

383 Distribution and frequency of mite species are given in Supplementary Material Table 7. A total of 40 384 water-mite taxa were observed, 36 of which are members of the true water mites. With regard to 385 individual numbers, of the globally 827 specimens collected in the frame of this project, only 47 (5%), are 386 representative of terrestrial groups. They represented (specimen numbers in brackets): Trombidiformes 387 (3), Gamasida (4), and Oribatida (40, including 20 Phthiracaridae). True aquatic mites occurred in 14 of the 388 springs studied, but in six of these occurred in low numbers. Terrestrial taxa were recorded only from sites 389 GeRi MiSp-Su (Gamasida, Oribatida, 10 specimens) and FoVR PS-Sh (Oribatida, 1). Sites with very low 390 water-mite density were Bari_ShFS-mc (Partnunia aprutina, Protzia squamosa, each one specimen), Poia_MiSp-SC (Partnunia aprutina, one specimen), Laba_LPS-le (Aturus cf. natangensis, one specimen, 391 Sperchonopsis verrucosa, 2 specimens), FoVR_PS-Su (only one undetermined larva), FoVe_LCFS (Feltria 392 393 setigera, Lebertia schechteli, each 2 specimens), and CaLS LCFS (Pseudofeltria aemiliana, Protzia eximia gr., 394 1 specimen each). Furthermore, also CiLi HygS (3 water mite taxa, 14 specimens) and Cara LPS-sn (4 water 395 mite taxa, 5 specimens) had a rather poor fauna. Higher population densities and species richness were 396 observed at sites MtPe_ShFS-Hi, MaPa_ShFS-Hi, Laga_ShFS-Hi, MtNe_ShFS-Hi, Prin_ShFS-pH and 397 Helo_LCSe.

398 Hygrobates psammocrenicus occurred in greatest abundance, mostly due to an extreme density of this 399 species at one site: Laga ShFS-Hi. Sperchon thienemanni was the next most common taxon, along with 400 Feltria setigera and Partnunia aprutina, as frequently recorded species (4 sites). Other species found in 401 higher numbers or more frequently (specimen numbers/frequency in parentheses) are Lebertia schechteli 402 (36/3), Atractides longisetus (24/2), Pseudofeltria appenninica (14/2), Hydrovolzia placophora (13/2), 403 Atractides loricatus (10/2), Sperchonopsis verrucosa (7/3), and Panisus michaeli (4/3). The remaining 25 404 species were found at one or two site(s) only and in low numbers, 14 of them as single specimens. 405 In this study, nearly 75% of the water mite species are classified to have a close relationship to spring 406 habitats: crenobionts (= strictly bound to springs): 60%; and crenophiles (= with a preference for springs,

but also in other habitats): 14%. Typical crenobionts, some of them with a particular preference for special
habitats, are *Panisus michaeli*, *Protzia squamosa paucipora*, *Sperchon resupinus*, *Bandakia concreta* (all
with a known preference for helocrenes), *Hydrovolzia placophora*, *Lebertia cuneifera*, *Lebertia holsatica*(preferably in rheohelocrenes rich in macrophytes and fine detritus), and *Lebertia schechteli*, *Sperchon mutilus*, *Sperchon thienemanni* (in all types of springs). Important crenophiles in our study are *Lebertia maculosa*, *Sperchon squamosus* and *Atractides loricatus*.

The pattern revealed by y-diversity partitioning (Table 3) parallels the results obtained for diatoms.

414

415

416 4. Discussion

The regional assemblages of spring organisms examined included a very high total number of species,
including several new or putatively-new species, and many Red-List taxa. The γ-diversity partitioning for
both diatoms and water mites demonstrated that α-diversity of the individual springs is relatively low, and
that the main contribution to γ-diversity was due to β-diversity, that is, a high turnover of species among

421 springs. This pattern and the percentages were very similar for diatoms and water mites.

422 These results are in good agreement with previous studies. Nascimbene et al. (2011) investigated algae,

423 diatoms, lichens, and bryophytes in springs of the southeastern Alps and compared α -, β - and γ -diversity: In

424 individual springs, these photoautotroph groups formed relatively species poor communities with a high

425 species replacement among springs whilst regional species pools were important. Studying

426 macrozoobenthos from 19 mountain springs of the River Sarca catchment in the Adamello-Brenta Nature

427 Park, Bonettini and Cantonati (1996), observed low species numbers in individual biotopes, and a high total

428 number of taxa. This condition demonstrates marked site to site (spatial) differences, which also emerged

429 from the TWINSPAN ordination. In the present study, springs with the greatest number of species, and

those that may be considered rare in northern Italy, were characterized by low flow variability. Pascual et

431 al. (2020) underline the uniqueness of individual spring communities after studying α -, β - and γ -diversity of

macroalgae, diatoms, bryophytes, vascular plants, aquatic invertebrates and vertebrates in Mediterranean
springs. Our findings also have important implications for conservation, suggesting that the protection of
single sites will not be effective, because a biodiversity conservation plan for spring environments requires
a plan developed at the regional level, and include a network of sites representative of the different
ecomorphologies, lithologies, and other key ecological factors.

These results also reveal the need for a deeper understanding of the causes of high γ-diversity.
Specifically, these data demonstrate that high geological diversity (lithotypes, aquifer hydrostructure, etc.),
translates into high spring ecomorphological and hydrochemical diversity. Further, these patterns are
reflected in high biodiversity. Other factors that may affect high γ-diversity include ecotonal character
(aquatic-aerial) of spring habitats, the fact that spring habitats in the study region (Emilia-Romagna) are still
relatively unexplored, and a general lack of high spatial resolution and in-depth taxonomic approaches of
springs.

444 A large part of the geological framework of the Northern Apennines comprises Cretaceous to Miocene, 445 marine neritic (shallow water) or turbiditic (deep water) deposits (siliciclastic arenites, marly-limestones or 446 arenaceous-pelitic turbidites), as well as scattered outcrops of ophiolites, mostly composed of peridotites, 447 serpentinites, gabbros, and basalts. The latter represent inducted remnants of the original oceanic crust of 448 the Ligurian basin, developed in the Middle to Upper Jurassic, which separated the European from the 449 Adriatic plate (Marroni et al., 2010). Each of these geological units are tectonically layered and deformed 450 within the Apennine system, and constitute valuable groundwater reservoirs. In some circumstances, the 451 sedimentary rocks exhibit heterogeneous and anisotropic hydraulic conductivity distributions, similar to 452 those commonly observed for HRA (Gargini et al., 2006; 2008; Vincenzi et al., 2009; Piccinini et al., 2013; 453 Vincenzi et al., 2014; Segadelli et al., 2017b). In particular, Gargini et al. (2014) and Piccinini et al. (2013) 454 consider that the Northern Apennines groundwater reservoirs are all represented either by hard rock 455 aquifers sensu stricto (like ophiolites) or by sedimentary units comprising hard rock aquifers as defined 456 above (Fig. 1). These units can be classified as shallow/surficial, recharge-dominated, strongly 457 heterogenous and anisotropic aquifers. Much less represented are the karstic aquifers, comprising Triassic -

458 Messinian evaporite sequences and Late Jurassic - Paleogene non-metamorphic carbonate successions (Fig. 459 1). The geological, structural and geomorphological survey conducted in part during the EBERs project and 460 in subsequent years (Gargini et al., 2014; Cantonati et al., 2016; Segadelli et al., 2017a;b; Cantonati et al., 2020) highlighted in the study area a variability of hydrogeological situations as described and summarized 461 462 in the conceptual scheme proposed in Figs 1-2, with particular attention to the HRA group, because it 463 represents, from a geological standpoint, the backbone of the northern Apennines. In one system, very 464 high pH values were associated with low temperature reaction between meteoric water and ultramafic 465 rocks (Neal and Shand, 2002; Boschetti and Toscani, 2008; Boschetti et al., 2013).

466 The percentage of Red List diatom species identified in this study belonging to the most important 467 threat categories (60%) is remarkably high. As a comparison, ~ 50% Red List species belonging to threat 468 categories were previously found in high-integrity springs and high-mountain lakes of the Alps (Cantonati et 469 al., 2012a). It is also important to note that most species belonging to the highest threat categories 470 encountered were collected in low-conductivity and dystrophic mountain springs, habitats that were 471 singled out for their rich and peculiar diatom microflora by previous works (Cantonati et al., 2009; 2011; 472 2019), and, secondarily, in limestone-precipitating springs (LPS), which are as well known as special habitats 473 (Cantonati et al., 2016). The share of Red-List threat categories species was also high likely because in the 474 recently published Red List (Hofmann et al., 2018; summarized in Supplementary Material Table 3), the 475 status of most species changed as a result of the new Red List classifications (Lange-Bertalot, 1996), with 476 many placed in higher threat categories. The relatively low share of aerial diatom species in the present 477 study (less than 10%) is likely due to a focus on stable-discharge springs for this study, as compared to 478 previous investigations in the Alps. Most diatom, bryophyte and vascular-plant species were encountered 479 only once or twice because of the high diversity of spring typologies and the consequent distinctiveness of 480 the individual spring sites.

In the zoobenthos, a considerable abundance and frequency of stygophilous taxa were observed, which suggests that the selected sites are probably of considerable biogeographical interest. As confirmed also by this study, in Europe, but probably also on other continents, water mites (Hydrachnidia) are the group of

484 organisms (other than microbes and diatoms) with the greatest number and percentage of species, which in many different genera, independently evolved strong ecological links to spring habitats (Gerecke et al., 485 486 2018). Only in Hydrobiid snails, a similarly high share of spring-typical species is reported from western North America, due to a strong radiation of the genus Pyrgulopsis (Hershler et al., 2014). As with diatoms 487 488 (Hofmann et al., 2018), these data offer information useful for a future Red List for European water mites. 489 In the course of the EBERs study, *Pseudofeltria aemiliana* was detected and described as a species new to 490 science (Gerecke, 2014). Unusual in water mites, and particularly surprising in a genus with strong sexual 491 dimorphism, this species is more distinct in females than in males. As with all known species of the genus, it 492 probably is a crenobiont with a preference for weakly seeping helocrenes. It can be considered an endemic 493 species of the Northern Apennines. The material from the study area includes also further interesting 494 records: the species found in highest abundance, *Hygrobates psammocrenicus*, has been described only 495 recently from springs in the Apennines and on Corsica (Gerecke and Di Sabatino, 2013); Stygothrombium 496 chappuisi is a very rare species reported from Italy only recently (Veneto, Emilia Romagna: Gerecke and Di 497 Sabatino, 2013); and Partnunia aprutina, after its first description from Abruzzo (Gerecke, 1993), has been 498 found again recently in several sites of the northern Appennines (Bottazzi et al., 2011). Lebertia fontana in 499 Italy was previously known only from two sites in Trentino (Gerecke, 2009), so this is the first record South 500 of the Alps, while Lebertia mediterranea, described from several sites in Italy, France (Corsica), Spain and 501 Bulgaria, and later on detected also in the Austrian Alps (Fišer et al., 2012), is a species in need of revision. 502 Specimens of the latter species found in this study differ from the original diagnosis in some morphological 503 details. With reference to species closely linked to groundwater habitats, the most striking result was the 504 finding of a large population of the typical interstitial-dwelling, laterally-compressed water mite 505 Frontipodopsis reticulatifrons. This species is normally restricted to interstitial waters, and not found in 506 springheads.

507 The topic of the relationship between biodiversity and geodiversity is now emerging with greater 508 awareness in the international scientific community (Kärnä et al, 2019; Alahuhta et al., 2020). There is a 509 clear need to develop new approaches through which to quantify and allow the comparison between

geodiversity and biodiversity in many types of ecosystems. In particular, as is already the case in other
ecological questions, the adoption of suitable indices and metrics stimulates and facilitates the comparison
between the biotic and abiotic diversity of a given area.

513Our data suggest that only an integrated hydrogeological-ecological approach (= ecohydrogeological514perspective; Cantonati et al., 2020) will permit researchers to accurately define the foundations for515conservation actions and for the monitoring of springs. These systems should not be examined as simple516points of aquifer-system discharge, but as ecotones with regional patterns that create complex GDEs517(groundwater dependent ecosystem, Bertrand et al., 2012). Springs, studied in an ecohydrogeological518perspective, are ideal systems in which to investigate and understand the geo-biodiversity relationship.

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- 520

521 **5. Conclusions**

522 The main conclusions that can be drawn from our observations are as follows:

- We confirm for spring habitats that the study region (Emilia-Romagna) is highly complex geoscape
 and an area of high geodiversity. In relation to inland waters, and spring ecosystems in particular,
 this is mainly apparent in the occurrence of diverse lithotypes and aquifer structures.
- Our data demonstrate that the diversity of rock and aquifer types generates a wide variation in
 ecomorphological types and hydrochemistry.

Considering two groups of organisms (diatoms, water mites) with contrasting ecological roles
 within the spring ecosystem, our study for a deeper understanding of spring-habitat biodiversity
 and uniqueness and of their determinants shows that the multiplicity of ecomorphological and
 hydrochemical types translates into high landscape-level (γ) biodiversity, with individual sites
 typically having relatively low diversity (α diversity) but differing markedly from one another (high
 individuality).

534	• The main consequence of our findings for spring-ecosystem conservation is that it is imperative to
535	protect representative and proportional groups of springs at the landscape level.
536	We further note that springs, studied from an ecohydrogeological perspective, are ideal systems where
537	to investigate geo-biodiversity relationships.
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539	
540	Acknowledgements
541	We are grateful to the Geological Survey of the Emilia-Romagna Region (in particular Raffaele Pignone
542	and Gabriele Bartolini) for fostering and funding the EBERs (Exploring the Biodiversity of Emilia-Romagna
543	springs; 2011-2013) Project, which provided all data used in this paper. Many thanks go to Horst Lange-
544	Bertalot for support in the identification of complex diatom taxa.
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751 Figure legends

753	Fig. 1. Geological map with the location of the springs studied. The small images exemplify the geodiversity
754	that characterizes the landscape of the Emilia-Romagna Region. (4) The basalts of Mt. Ciapa Liscia, 1658 m
755	a.s.l. (9) The serpentinized peridotites of Mt. Prinzera, 736 m a.s.l. (1) View to the East from the Fugicchia
756	Pass: siliciclastic arenites, 1667 m a.s.l. (12) Parallel-stratified calcarenite deposits, 636 m a.s.l. See Table 1
757	for the meaning of the codes of the 16 springs.
758	
759	Fig. 2. Hydrogeological block diagram of groundwater flow systems in the main hard rock aquifers
760	outcropping in the study area. Spring numbers are as in Fig. 1.
761	
762	Fig. 3. Representative images of springs of the different ecomorphological types and subcategories.
763	
764	Fig. 4. Box plots of the main environmental biota determinants.
765	
766	Fig. 5. LM and SEM (3a external view; 9e internal view) micrographs of selected frequent and abundant,
767	characteristic, rare and Red List diatom species of the different ecomorphological types. The micrographs
768	(sites) follow the usual order of increasing mineralization. 1a-d, Helo_LCSe, 1a Navicula angusta, 1b
769	Gomphonema pseudobohemicum, 1c Eunotia soleirolii "dwarf", 1d Stauroneis smithii; 2a-d, CaLS_LCFS, 2a
770	Encyonema neomesianum, 2b Eunotia cisalpina, 2c Chamaepinnularia muscicola, 2d Kolbesia carissima; 3a -
771	d, FoVe_LCFS, 3a Geissleria acceptata, 3b Gomphonema sphenovertex, 3c Nitzschia dealpina, 3d
772	Achnanthidium lineare; 4a-c, CiLi_HygS), 4a Gomphonema minusculum, 4b Encyonema ventricosum, 4c
773	Amphora micra; 5a-c, MtNe_ShFS-Hi , 5a Encyonema vulgare, 5b Encyonema sublangebertalotii, 5c
774	Encyonopsis moseri; 6a, MtPe_ShFS-Hi, 6a Brachysira calcicola; 7a-c, Laga_ShFS-Hi, 7a Sellaphora
775	atomoides, 7b Sellaphora seminulum, 7c Gomphosphenia fontinalis girdle view; 8a-b, MaPa_ShFS-Hi), 8a

776 Navicula splendicula, 8b Neidiomorpha binodiformis. 9a-f, Prin_ShFS-pH, 9a Gomphonema auritum, 9b Navicula veronensis, 9c-d Humidophila irata, 9e Reimeria uniseriata, 9f Encyonema bipartitum; 10a, 777 778 Bari_ShFS-mc, 10a Gomphonema angustum; 11a-c, Cara_LPS-sn, 11a Gomphonema lateripunctatum, 11b 779 Achnanthidium trinode, 11c Gomphonema tenoccultum; **12a-d**, **Laba_LPS-le**, 12a Cymbella tridentina, 12b 780 C. tumidula, 12c-d Denticula elegans: 12c valve view, 12d girdle view; 13a, FoVR_PS-Su, 13a Fragilaria 781 famelica low-salinity morphotype; 14a-c, FoVR_PS-Sh, 14a Staurophora wislouchii, 14b-c Planothidium 782 werumianum: 14b RL valve, 14c raphe valve; 15a-b, Poia_MiSp-SC, 15a Nitzschia frustulum, 15b Fragilaria 783 famelica.

784

785 Fig. 6. Site-environmental variables, and species-environmental variables CCA biplots for the epilithon and 786 the epibryon, respectively. See Table 1 for the meaning of the codes of the 16 springs. Species OMNIDIA 787 acronyms: PDAO = Psammothidium daonense, ADOL = Achnanthidium sp. aff. dolomiticum, ADTR = 788 Achnanthidium trinode, NULA = Nupela lapidosa, GTNO = Gomphonema tenoccultum, ONEO = Odontidium 789 neolongissimum, GLAT = Gomphonema lateripunctatum, PMCR = Psammothidium microscopicum, GCUN = 790 Gomphonema cuneolus, AINA = Amphora inariensis, NEXI = Navicula exilis, ADFO_mn = Achnanthidium 791 fontisalinae sp. nov. mn, AIND = Amphora indistincta, CRUM = Crenotia rumrichorum, COPL = Cocconeis 792 pseudolineata, FFAM = Fragilaria famelica, PTEN mn = Planothidium tenuilanceolatum sp. nov. mn; ACAF = 793 Achnanthidium affine, ENCM = Encyonopsis microcephala, DLMI = Delicata minuta, GPUM = Gomphonema 794 pumilum, PGRI = Psammothidium grischunum, GACC = Geissleria acceptata, HPEP = Humidophila perpusilla, 795 HUCO = Humidophila contenta, SENI = Sellaphora nigri, SSEM = Sellaphora seminulum. 796

Fig. 7. Scatterplots showing the relations between diatom-epibryon Shannon-Wiener diversity (*H'*) and
vegetation Shannon-Wiener diversity (*H'*) and lithology. Explanations of the eight values of the variable
'lithology' are given in Table 1.

- **Fig. 8.** Scatterplots showing the relations between selected zoobenthos groups (absolute abundances) and
- 802 lithology. The meaning of the eight values of the variable 'lithology' are explained in Table 1.

804	Supplemental online materials
805	
806	
807	Supplementary material Table 1. The seventy morphological, physical, and chemical variables and
808	parameters measured in the 16 springs.
809	
810	Supplementary material Table 2. Enrichment factors for the trace elements and metals measured in the
811	springs studied (2011).
812	
813	Supplementary material Table 3. List of all diatom species identified in the samples. N. occurrences, min,
814	max relative abundance %); Red-List status, aerial species, and ecology (according to Hofmann et al., 2018).
815	
816	Supplementary material Table 4. Bryophytes.
817	
818	Supplementary material Table 5. Vascular plants.
819	
820	Supplementary material Table 6. Zoobenthos.
821	
822	Supplementary material Table 7. Water mites.

Table 1. Ecomorphological and hydrogeological classifications of the springs studied and related variables.

Spring code	Helo_LC Se	CaLS_LCFS	FoVe_LC FS	CiLi_H ygS	MtNe_S hFS-Hi	MtPe_S hFS-Hi	Laga_Sh FS-Hi	MaPa_S hFS-Hi	Prin_Sh FS-pH	Bari_ShFS -mc	Cara_LPS -sn	Laba_LP S -le	FoVR_ PS-Su	FoVR_ PS-Sh	Poia_ MiSp- SC	GeRi_ MiSp- Su
Spring name	Helocre ne at Lago Scuro (Helo)	Capanne Lago Scuro (CaLS)	Fontana del Vescovo (FoVe)	Ciapa Liscia (CiLi)	Monte Nero (MtNe)	Monte Penna (MtPe)	Lagacci (Laga)	Mangia pane (MaPa)	Prinzera (Prin)	Barigazzo (Bari)	Carameto (Cara)	Labante (Laba)	Fonta nile Valle Re (FoVR) Sun	Fonta nile Valle Re (FoVR) Shade	Poiano (Poia)	Gessi Riolo (GeRi)
Ecomorphology	Low Cond. (LC) Seepage (Se)	Low Conduc Flowing Sp	ctivity (LC) rings (FS)	Hygro petric (Hyg) spring (S)	Shaded (Sh) Flowing Springs (FS) with Hildenbrandia (Hi)			ShFS with high pH (pH)	ShFS with medium cond. (mc)	LPS small near- natural (sn)	LPS large exploite d (le)	Pool Spring (PS) Sun (Su)	Pool Spring (PS) Shade (Sh)	Miner al Spr. (MiSp) Sodiu m Chlori de (SC)	Miner al Spr. (MiSp) Sulpha tes (Su)	
Cond . (uS cm ^{−1})	13	27	57	62	56	62	111	265	255	305	462	451	788	815	(3C) 11560	2170
Shading (%)	3	20	3	70	78	69	77	77	78	67	11	8	6	72	80	96
Discharge (L s ⁻¹)	0,01	0,8	1	1	1,2	3,5	3,5	0,9	0,03	0,4	0,07	15,5	0,5	0,5	50	2,5
рН	7,27	6,41	7,31	7,30	7,63	6,56	7,47	7,70	11,20	7,48	7,66	8,20	7,11	7,22	7,44	7,83
Na⁺ (mg L⁻¹)	1,5	1,3	1,6	1,5	0,39	1,8	1,0	2,8	33	6,7	4,4	4,8	21	22	2250	11,1
Cl ⁻ (mg L ⁻¹)	1,7	1,2	1,1	1,4	0,7	1,2	1,0	1,5	24,0	2,7	2,6	7,5	41,0	41,0	4370,0	13,6
SO 4 ²⁻ (mg L ⁻¹)	1,7	1,5	1,8	2,8	0,81	2,5	2,2	10,7	9,7	27	45	27	69	68	2090	144
Lithology var. (1-8)	1	1	1	2	3	2	3	4	4	5	6	6	7	7	8	8
Lithology	silio	ciclastic areni	ites	basalts peridot basalts es peridot. lithic lin					limest. flyschs; calcarenites		gravel a	nd sand	gyp	sum		
Aquifer		turbidites		ophiolites turbidites							alluvial deposits		evaporites			
Aquifer broad cat.					I	Hard Rock	Aquifers (H	IRA)					Porous	aquifer	Ка	rst

Coving code	Helo_	CaLS_	FoVe_	CiLi_	MtNe_	MtPe_	Laga_	MaPa_	Prin_	Bari_	Cara_	Laba_	FoVR_	FoVR_	Poia_	GeRi_
Spring code	LCSe	LCFS	LCFS	HygS	ShFS-Hi	ShFS-Hi	ShFS-Hi	ShFS-Hi	ShFS-pH	ShFS-mc	LPS-sn	LPS-le	PS-Su	PS-Sh	MiSp-SC	MiSp-Su
Longitude E	10°2'56.534"	10°2'54.256"	10°2'26.790"	9°28'2.709"	9°31'13.674"	9°30'29.493"	9°35'41.766"	9°29'24.606"	10°4'45.008"	9°48'37.719"	9°45'35.165"	11°2'9.836"	10°31'22.143"	10°31'20.988"	10°26'21.274"	11°43'50.073"
Latitude N	44°22'41.151"	44°22'40.381"	44°22'44.767"	44°34'14.370"	44°33'38.595"	44°29'6.029"	44°37'53.091"	44°46'19.328"	44°38'45.603"	44°36'44.062"	44°39'50.840"	44°15'38.339"	44°45'53.362"	44°45'50.898"	44°23'20.168"	44°14'8.570"
Elevation (m a.s.l.)	1534	1547,5	1613	1381	1509	1324	926	736	492	880	758	603	31,3	32	430	167,5
Discharge (L s ⁻¹)	0,01	0,8	1	1	1,2	3,5	3,5	0,9	0,03	0,4	0,07	15,5	0,5	0,5	50	2,5
Variab. Ind. Disch.	200	50	269	172	98	70	162	53	95	93	150	233	20	20	129	200
T (°C)	9,8	8,8	4,7	5,4	5,8	5,3	7,7	11,1	14	7,9	12,8	15,4	15	13,2	9,6	11,9
Conduct . (μS cm⁻¹)	13	27	57	62	56	62	111	265	255	305	462	451	788	815	11560	2170
рН	7,27	6,41	7,31	7,3	7,63	6,56	7,47	7,7	11,2	7,48	7,66	8,2	7,11	7,22	7,44	7,83
HCO3 ⁻ (mg L ⁻¹)	14	8,2	18	20	20	19	39	93	40	92	145	143	210	208	79	110
Ca²⁺ (mg L ⁻¹)	6,9	4,2	10,7	11,5	0,77	10,3	2,4	23	10,3	43	88	69	143	142	880	651
Mg²⁺ (mg L ⁻¹)	0,63	0,42	0,53	1,7	7,3	1,5	14,7	25	0,11	15	15,5	23	29	29	4,1	39
ΤΡ (μg L ⁻¹)	<1	4	14	3	<1	7	6	<1	10	<1	6	4	43	3	33	11
NO 3 ⁻ (mg L ⁻¹)	0,17	1,1	1,1	0,43	0,46	1,2	0,64	1,1	<0.05	1,1	0,12	1,9	41	48	0,56	14,6
Cr (µg L⁻¹)	0,28	0,24	0,5	1,1	40	0,88	16,8	15,4	0,18	0,85	0,12	0,38	1	0,29	3,4	1,06
Ni (μg L ^{⁻1})	0,54	0,15	0,19	0,31	2,6	0,24	4,4	10,8	0,14	0,74	0,74	1,33	3,3	1,35	3	5,4
Cu (µg L ^{⁻1})	1,11	0,24	0,13	0,11	0,17	0,17	0,14	0,93	0,54	0,18	0,19	0,17	1,11	0,25	7,1	0,33
Zn (µg L ^{⁻1})	1,21	0,36	0,2	0,13	1,15	0,62	0,32	0,71	0,28	0,26	0,36	0,2	3	0,44	0,32	0,38
Fe (µg L ^{⁻1})	15	9,3	18	28	7,2	14	13	82	8,4	59	58	38	391	93	274	643
Mn (μg L ^{⁻1})	11,1	0,15	0,061	0,079	0,26	0,079	0,18	1,9	0,26	1,6	0,33	0,078	10,2	0,43	0,52	0,23

Table 2. Main morphological, physical, and chemical characteristics as emerging from data processing and statistical analyses.

S	epilithon	epibryon	debris	Η'	epilithon	epibryon	debris
min	7	7	5	min	0,56	0,25	0,52
max	46	65	22	max	2,59	2,15	2,29
mean	25	28	16	mean	1,74	1,57	1,41
N	12	14	3	Ν	12	14	2

Table 3. Diatom species richness (S) and Shannon-Wiener diversity (H').

epilithon	df	Chi2	F	Р	epibryon	df	Chi2	F	Р
elevation	1	0,575	1,343	0,039	elevation	1	0,636	1,413	0,029
logNO3	1	0,523	1,222	0,132	НСОЗ	1	0,540	1,200	0,228
Cr	1	0,657	1,535	0,038	Mg	1	0,418	0,930	0,562
					logNO3	1	0,726	1,613	0,044
Residual	10	4,280			Residual	9	4,050		

Table 4. Diatom (epilithon and epibryon) Canonical Correspondence Analysis (CCA) data.

Table 5. Landscape-level (γ) diversity partitioning for diatoms and water mites.

	DIATOMS												
		ness	S	hannon		Richness							
	epilith	on	epibry	on	epilith	on	epibry	on					
	statistic	%	statistic	%	statistic	%	statistic	%	statistic	%			
alpha.1	23,5	0,12	28,4	0,13	1,69	0,51	1,57	0,50	5,07	0,13			
gamma	199		214		3,35		3,13		39				
beta.1	175,5	0,88	185,6	0,87	1,65	0,49	1,56	0,50	33,9	0,87			















