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4 **Groups of small lakes maintain larger microalgal diversity than large ones**

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20

21 **Abstract**

22 The question of whether one large, continuous area, or many smaller habitats maintain more
23 species is one of the most relevant questions in conservation ecology and it is referred to as
24 SLOSS (Single Large Or Several Small) dilemma in the literature. This question has not yet
25 been raised in the case of microscopic organisms, therefore we investigated whether the
26 SLOSS dilemma could apply or not to phytoplankton and benthic diatom metacommunities.
27 Benthic diatom and phytoplankton diversity in pools and ponds of different sizes (ranging
28 between 10^{-2} - 10^7 m²) was studied. Species richness of water bodies belonging the
29 neighbouring size categories was compared step by step across the whole size gradient. With
30 the exception of the compared 10^4 – 10^5 m² and 10^5 – 10^6 m² size categories, where
31 phytoplankton and benthic diatom richness values of the SL water bodies were higher than
32 that of the SS ones, diversity of several smaller (SS) sized waters was higher than that in
33 single large ones (SL) throughout the whole studied size range. The rate of the various
34 functional groups of algae, including both the benthic diatoms and phytoplankton, showed
35 remarkable changes from the smaller water bodies to large sized ones.

36 Keywords: SLOSS-dilemma, lakes, benthic diatom, phytoplankton, wide size scale

37 **1. Introduction**

38 The question of how cumulative species richness in several small habitats relates to that in
39 one large area (where cumulative area of SS is equivalent to that of SL) became known as the
40 SLOSS-debate (Single Large Or Several Small) in ecology. Several studies on the SLOSS
41 dilemma were triggered by the frightening rate of habitat fragmentations which became an
42 important issue in nature conservation (Foley et al., 2005). Since understanding the SLOSS-
43 dilemma may help to find the optimal size of nature reserves it has been studied for decades
44 by many authors since the seventies (Diamond, 1975; Wilson and Willis, 1975; Simberloff
45 and Abele, 1976). While many studies demonstrated, that from the conservational point of
46 view, several small habitats can be as valuable as a single larger-sized one (Turner and
47 Corlett, 1996; Honnay et al., 1999; Gibb and Hochuli, 2002), there are many opposing results
48 in the literature, which stress the importance of a single large habitat (Matias et al., 2010; Le
49 Roux et al., 2015). The contradictory findings of these studies indicate that this debate is still
50 unresolved (Tjørve, 2010; Rösch et al., 2015).

51 The size of the suitable habitat is largely determined by the characteristics of the species,
52 which tries to settle and establish residence. Those species that are typically generalists or
53 opportunists can easily adapt to the conditions of different-sized habitats (Gibb and Hochuli,
54 2002). High dispersal capability, that is characteristic for birds, allows them to survive in
55 small habitats in the same way as in larger ones (Lindenmayer et al., 2015). On the other
56 hand, the single large habitat ensures appropriate conditions by minimizing the extinction rate
57 (Gaz and Garcia-Boyero, 1996; Le Roux et al., 2015). Besides the specific characteristics of
58 the studied taxa, contradictory findings can also be traced back to statistical uncertainties.
59 Theoretically, the SLOSS debate is in close connection with the species-area relationship
60 (SAR). Essence of the SAR's theory is that the species richness increases with the increasing
61 area size. This relation has been demonstrated for various organisms both on macro- (Connor

62 and McCoy, 1979; Tjørve, 2003; Báldi, 2008; Lindenmayer et al., 2015; Matthews et al.,
63 2016) and micro-scale (Smith et al., 2005; Bolgovics et al., 2016) and now, the SAR has
64 become an accepted conceptual framework for ecological researches. Besides its theoretical
65 importance, the species-area relationship (SAR) has substantial relevance from a nature
66 conservation point of view. Although on a large spatial scale SAR can be described well by
67 power function (Arrhenius, 1921), it becomes stochastic when only a small part of the size-
68 scale is studied. It is especially true for the lower end of the size scale, where, because of the
69 so called Small Island Effect (SIE) (Triantis and Sfenthourakis, 2011; Gao and Perry, 2016),
70 diversity changes in an unpredictable way.

71 Moreover, species-area relationship can also be interpreted within the framework of the
72 metacommunity theory (Gilpin and Hanski, 1991). This theory argues that local communities
73 are linked by dispersal of many potentially interactive species, and thus create a
74 metacommunity (Leibold et al., 2004). It means that, besides the local constraints, regional
75 processes (e.g. dispersal) have pronounced influence on the composition of local
76 communities. The most common distributional patterns in meta-communities are nestedness
77 and species turnover (Baselga, 2010). Nestedness means that within a metacommunity,
78 species of some local communities are the subsets of the larger, species rich communities;
79 while species turnover is the rate of species replacement in communities, which is a reflection
80 of habitat heterogeneity (Wiens, 1974; Astorga et al., 2014). These mechanisms shape the β -
81 diversity of communities (Harrison et al., 1992), which, however, can be partitioned by the
82 appropriate statistical tools (Baselga, 2010).

83 Majority of the above mentioned findings were obtained from studies on macroscopic taxa,
84 but investigations of the SAR or the SLOSS debate on microscopic organisms may have
85 similar relevance for the understanding of the compositional structure and functioning of
86 microbial ecosystems. Diverse microbial primary producer communities in the pelagic and

87 benthic zone sustain diverse grazer assemblages, have an impact on their composition and
88 growth rate, and have far-reaching consequences for the structure and functioning of the
89 whole aquatic food web (Liess and Hillebrand, 2004; Striebel et al., 2012).

90 Lakes and ponds are ideal objects to investigate the SLOSS-dilemma across a large spatial
91 scale, because they can be considered as aquatic islands on a terrestrial landscape and their
92 size range may cover several orders of magnitude even within a small geographic area
93 (Dodson, 1992). These habitats provide suitable conditions for various aquatic organisms
94 from the microscopic to the macroscopic ones. Among these organisms, algae represent a
95 group which is usually characterized by high species richness and consists of taxa that are
96 relatively easy to identify. These attributes make them suitable to answer various ecologically
97 relevant questions (Soininen et al., 2016; Török et al., 2016; Várbiro et al., 2017). In the last
98 decades, functional approaches were increasingly used in algal researches (Reynolds et al.,
99 2002; Padisák et al., 2009; Rimet and Bouchez, 2012; B-Béres et al., 2016, 2017; Tapolczai et
100 al., 2016). They can provide detailed information about the ecosystem functioning and ensure
101 a deep knowledge about ecosystem vitality. Thus, they have a remarkable role in conservation
102 and environmental management (Padisák et al., 2006; Borics et al., 2007; B-Béres et al.,
103 2019). In phytoplankton ecology, the functional group concept, proposed by Reynolds et al.
104 (2002), has become the most widely used classification system (Salmaso et al., 2015). Here,
105 algae and cyanobacteria are classified into more than 40 FGs based on their habitat
106 preferences and environmental tolerances (Padisák et al., 2009; Salmaso et al., 2015). In
107 diatom ecology, the use of functional classifications is based on morphological, behavioral
108 and physiological criteria (Passy, 2007; Rimet and Bouchez, 2012; Berthon et al., 2011).
109 Merging these approaches enabled the establishment of 20 combined eco-morphological
110 functional groups (CEMFGs) by B-Béres et al. (2016). The feasibility and utility of this
111 system have been studied under different environmental conditions (lowland rivers and

112 streams - B-Béres et al., 2017; continental saline lakes and ponds - Stenger-Kovács et al.,
113 2018).

114 While the relationship between nutrients and phytoplankton biomass has been well
115 demonstrated, nutrient-diversity relationships might potentially exist only in oligotrophic or
116 oligo-mesotrophic range (Soininen and Meier, 2014), where the low nutrient concentration
117 might act as an environmental filter. In nutrient- enriched aquatic environments, causal
118 relationship between nutrient availability and species richness could not be proved (Várbíró et
119 al., 2017). In these systems the number of within-lake microhabitats has pronounced influence
120 on species diversity (Görgényi et al., 2019). Eutrophic lakes of the Carpathian Basin therefore
121 are appropriate objects to study the size-related aspects of diversity. Studying the SLOSS
122 debate on microbial aquatic organisms is not just a theoretical issue but it might also have
123 conservational relevance. In this study, we have performed an extensive analysis of the
124 SLOSS debate on a large spatial scale in Hungary using both benthic diatoms and
125 phytoplankton.

126 We addressed the following hypotheses:

- 127 (i) since we expect higher complexity in the larger water body categories, species
128 richness of single large (SL) water bodies will be higher than species richness of
129 several small (SS) ones
- 130 (ii) in accordance with the small island effect (SIE) species richness in smaller size
131 categories (10^{-2} - 10^2 m²) will change randomly, and clear patterns in the SLOSS
132 dilemma will not be observed,
- 133 (iii) since increasing complexity is expected with the increasing habitat size, this
134 complexity will result in higher number of functional groups in the case of both
135 studied group.

136

137 **2. Material and methods**

138 **2.1 Study area**

139 Testing the research hypotheses eutrophic pools, ponds and lakes of varying sizes were
140 selected in the whole area of Hungary (Central Europe). The area of the studied lakes covered
141 10 orders of magnitude, extending from 10^{-2} to 10^7 m².

142 The data are partly derived from the National Hungarian Database, which contains
143 phytoplankton and phytobenthon data for shallow lakes (mean depth <3m) and ponds between
144 10^3 - 10^7 m² areas. To acquire the surface area of these ponds, oxbows and other larger standing
145 water bodies we used the data of the national Hungarian database (database 1).

146 Samples belonging to the five smaller size categories (10^{-2} - 10^2 m²) were collected from an
147 extended area that was used as a bombing and gunnery training range between 1940 and 1990
148 and later for pasturing. This area is situated in the Hungarian Great Plain (Hungary, 47° 27'
149 00.36" N and 20° 59' 44.09"), and the intensive bombing created thousands of bomb crater
150 ponds of different sizes (10^0 - 10^2 m²) during the decades. In this area, very small pools were
151 also created by grazing of the animals. Their sizes varied from 10^{-1} to 10^{-2} m². To calculate the
152 area of the small pools (10^{-2} - 10^2 m²) at the bombing range we measured their linear
153 dimensions by a tape measure. Limnological characteristics of studied lakes can be seen in
154 Table A.1.

155 **2.2 Sampling and sample processing**

156 **2.2.1 Diatoms**

157 The sampling and sample processing of benthic diatoms were done according to international
158 standards (EN 13946, EN 14407). From shallow lakes and ponds with 10^3 - 10^7 m² area, and
159 from the bomb crater ponds with 10^0 - 10^2 m² area samples were collected from reed stems. At
160 those sites where macrophytes were unavailable (10^{-2} – 10^{-1} m² size range), samples were
161 taken from the psammon. Although differences in substrata types might cause differences in

162 the relative abundance of the occurring elements but the species composition of psammon to
163 the harder substrates is similar (Townsend and Gell, 2005). Similar results were found by
164 Szabó et al. (2018) studying the benthic diatom flora of lakes and ponds in Hungary: They
165 found no significant differences in the composition and diversity of algal assemblages
166 collected from different substrates.

167 Samples from shallow lakes and ponds ($10^3 - 10^7$ m² size range) were collected in the
168 growing season between 2001 and 2012, while samples from small ponds in the bombing
169 range were taken in September 2011.

170 In order to make the diatom valves clearly visible in benthic samples, 2 cm³ H₂O₂ were added
171 to 1 cm³ sample. In addition, a few drops of HCl were also added to remove calcium
172 carbonate. In the next step, the samples were placed in a water bath for one day at 70 °C.
173 Finally, permanent slides were made with Cargille-Meltmount mounting medium (refractive
174 index = 1.704). Diatom species were identified with Zeiss Axioimager A2 upright microscope
175 at 1000 × magnification. Additionally, oil immersion and differential interference contrast
176 (DIC) technique were applied. A minimum of 400 valves were counted per slides.

177

178 2.2.2 Phytoplankton

179 The sampling and sample processing of phytoplankton were done according to international
180 standards (EN 16698, EN 16695, EN 15204). In the case of smaller sized pools (10^2 - 10^3 m²)
181 phytoplankton samples were taken from the middle of the pools by a plastic dish in the second
182 half of the vegetation period 2011. In the case of the shallow lakes and ponds (10^3 - 10^7 m²)
183 samples were collected in the vegetation period between 2001 and 2012. In these water bodies
184 more sample sites were designated in the representative points of the lakes. Samples were
185 collected from the euphotic layer with tube sampler. The euphotic layer was considered as 2.5
186 times of the Secchi depth. These subsamples were mixed in a larger plastic container, from

187 which 0.5 L of water was taken and fixed with formaldehyde solution (concentration of 4%)
188 and stored in darkness at 4 °C.

189 Phytoplankton samples were settled in 5 ml sedimentation chambers for 24 hours, and then
190 analysed by inverted microscopes (Utermöhl, 1958), applying 400× magnification. To
191 estimate the relative abundance of smaller algal units a minimum of 400 specimens were
192 counted. The entire area of each chamber was investigated to estimate the number of large
193 sized taxa. The list of the studied lakes and the observed number of samples are shown in
194 Table 1.

195

196 **2.3 Area of the SL and SS lakes**

197 Since we hypothesised that the values of the metrics used for representing the SLOSS depend
198 on the size of the water bodies, all adjacent size categories were separately compared within
199 the studied size range (10^{-2} - 10^7 m²) (Fig. 1). More precisely it means, that taxonomical and
200 functional diversities of the smaller water body category were compared to metrics of waters
201 in the next larger category.

202 In an ideal case the sum of the area of small water bodies is equal with the area of the single
203 large one. However, our database did not make possible that the area of SS lakes would be
204 equal to that of the SL one. As it is illustrated in Fig. 2, in the majority of cases, the sum of
205 the area of the SS lakes was smaller.

206 Within this smaller size range (10^{-2} - 10^2 m²), where we had five pools in each size category,
207 the size of SL pools was twice as large as that of the SS pools. In the larger size categories
208 (10^3 - 10^7 m²) the area covered by the SS lakes also showed differences.

209

210 **2.4 Species richness estimations - ESR**

211 The observed number of species occasionally might give a biased estimate of the real species
212 richness, and the bias is mostly related to differences in the sampling effort, therefore one
213 major challenge in SLOSS studies is how to compare the species richness of the different
214 areas. Since in the smallest size categories (10^{-2} - 10^2 m²) single samples were collected from
215 every water body, in the case of these waters statistical richness estimations cannot be
216 applied. However, with respect to the small size of these water bodies, the sample volume/
217 habitat volume ratios were high, which increased the detectability of an individual algal unit.
218 Since higher individual detectability increases the detection of species (Buckland et al., 2011),
219 the observed number of species well represented the real species richness in these small
220 habitats. In these size categories richness values of the SS lakes were considered as the sum of
221 the observed species numbers of the 5 small pools. Species richness of the SL lake (i.e. lake in
222 one order of magnitude larger size category) was considered as the mean of the observed
223 richness values of the 5 pools belonging to the given category.

224 In the case of larger size categories (10^3 - 10^7 m²), data for longer time periods were available.
225 Although we had different numbers of samples from each lake in all size categories (Fig. 3A),
226 these sample numbers were sufficient to apply a more rigorous statistical comparison between
227 the richness of SL and SS lakes.

228 Since the species numbers increase with the number of the samples studied, our aim was that
229 in the pairwise comparisons between SL and SS lakes the number of samples considered
230 would be equal. To achieve this, we applied Chao's sample-based extrapolation technique
231 (Chao et al., 2014), which is a non-asymptotic approach, that enables us to compare diversity
232 estimates by using seamless rarefaction and extrapolation (R/E) sampling curves. In the case
233 of phytoplankton, the databases usually contain species specific biomass data, which do not
234 enable the application of individual-based rarefactions. However Chao's method is an

235 incidence-based technique, which considers the occurrences of species within the given
236 sample, but ignores relative abundances.

237 Increasing lake size means decreasing individual and species detectability, therefore parallel
238 with an increase in the lake size, we proposed to consider increasing sample numbers in
239 richness comparisons (Table 1). To estimate the richness in SL lakes (ESR_{SL}) using the
240 extrapolation curves, we calculated the species richness for the proposed sample numbers for
241 each lake in the given size category (Fig. 3C), and means of these values were considered as
242 ESR_{SL} values.

243 When estimating the species richness of SS lakes (ESR_{SS}), as a first step, species occurrence
244 matrices of all lakes within the given size category were stacked. In the next step, applying
245 the sample numbers that were considered for calculations of ESR_{SL} in the one order of
246 magnitude larger size category, we calculated estimated species richness of the SS lakes (Fig.
247 4C).

248 These procedures were repeated in the case of each pairwise comparison. Finally, to represent
249 the SLOSS dilemma, the quotient ESR_{SL}/ESR_{SS} was plotted against the area of water bodies
250 (Fig. 5).

251

252 **2.5 Evaluation of functional group richness and functional redundancy**

253 The observed differences between the functional group richness values of adjacent size
254 categories can be partly explained by functional differences between the compared water
255 bodies (see in subsection 2.3). These limnological and/or biological differences between water
256 bodies in adjacent size categories can result differences in the number of occurring functional
257 groups (FG) of benthic diatoms and phytoplankton (Table A.2 and A.3). Studying these
258 functional differences, taxa observed both in the benthic diatom and phytoplankton samples
259 were assigned to the appropriate FGs (Tables A.2 and A.3). Diatom species were assigned to

260 twenty combined eco-morphological functional groups according to B-Béres et al. (2016).
261 Functional classification of phytoplankton was based on the concept proposed by *sensu*
262 Reynolds et al. (2002); which was supplemented by Borics et al. (2007) and reviewed by
263 Padisák et al. (2009).

264

265 **2.6 Programs used for statistical analysis**

266 Rarefaction curves were drawn using the iNEXT (Hsieh et al. 2013, ver. 1.0) packages
267 available in R Studio (2012).

268

269 **3. Results**

270 Altogether 189 benthic diatom and 181 phytoplankton samples were collected from 36
271 different sized standing waters in Hungary. We identified 312 benthic diatom and 498
272 phytoplankton species in the samples.

273 The species richness of diatom assemblages in the SS lakes was higher at most size categories
274 (ESR_{SL}/ESR_{SS} values <1), except in the case of 10^5 m² size range (Fig. 6 A). At the 10^5 m²
275 size category more species could be observed in the SL lakes than in several smaller ones
276 (ESR_{SL}/ESR_{SS} value >1). The ESR_{SL}/ESR_{SS} values showed large variation in the small size
277 categories (from 10^{-2} m² to 10^2 m²), while they were more consistent in the case of larger
278 lakes (lake area $>10^3$ m²).

279 The results showed similar patterns in the case of the phytoplankton. The species richness of
280 SS lakes was higher in almost every size category, except in 10^4 m² area size (Fig. 6 B). The
281 values showed large variation across the whole size scale, but the data showed no discernible
282 trends or regularities. In contrast to benthic diatoms where ESR_{SL}/ESR_{SS} ratio showed only
283 small changes in the larger lake categories, phytoplankton richness of this lake size category

284 was considerably smaller than that in the sum of the lakes in the adjacent smaller lake size
285 category.

286

287 **3.1 Functional groups**

288 The number of functional groups showed similar patterns in the case of both benthic diatoms
289 and phytoplankton. Smaller values characterized the water bodies in the 10^{-2} m² to 10^2 m² size
290 range, while larger ones in the 10^3 - 10^7 m² range (Fig. 7 A-B, and Table A.2 and A.3).

291 Smaller differences could be observed in the larger lake categories where the number of
292 benthic diatom FGs was almost identical (~20), the phytoplankton FGs displayed a peak at
293 10^5 m² range and decreased thereafter.

294 The functional redundancies of benthic diatoms (i.e. number of species within the FGs)
295 showed characteristic changes along the size gradient (Fig. 8 A and Table A.2).

296 Richness of the motile groups decreased with water body size. An opposing tendency was
297 observed in the case of high profile groups which showed increasing redundancy from 10^3 m²
298 to the largest size categories.

299 The ratios of the phytoplankton functional groups also differed from each other in the case of
300 smaller and larger size categories (Fig. 8B and Table A.3).

301 In small sized water bodies (10^{-2} m² – 10^2 m²), the W1 functional group was dominant, that
302 mostly consists of euglenoid algae. In contrast to W1 group, richness of X1, N and Lo FGs
303 were higher in the larger size categories (for more information on functional groups see in
304 Table A.3).

305

306 **4. Discussion**

307 Our results clearly demonstrated that several small water bodies can maintain greater
308 phytoplankton and benthic diatom species richness than single large ones; thus the results did

309 not corroborate our first hypothesis. Considering that the aggregated areas of the several small
310 water bodies were smaller in almost each case of comparisons (Fig. 2), the results are even
311 more convincing.

312 In line with our second hypothesis the ESR_{SL}/ESR_{SS} values did not show any trends in the
313 case of small water bodies. Species numbers were lower and changed randomly in the smaller
314 size categories (10^{-2} - 10^2 m²) resulting in hectic changes in the ESR_{SL}/ESR_{SS} values. An
315 interesting interpretation of these results can be made in the context of the species-area
316 relationship (SAR). At large spatial scale, the SARs follow a power model (Arrhenius, 1921).
317 In contrast, the richness values change independently from the area in very small habitats,
318 resulting in unpredictable diversity patterns in these small habitats. This stochastic pattern has
319 been described as small island effect (SIE) in the literature of island biogeography (Lomolino
320 and Weiser, 2001; Triantis and Sfenthourakis, 2011). We think, that this phenomenon can
321 explain the large variations in the ESR_{SL}/ESR_{SS} ratio experienced in the case of small water
322 bodies.

323 Several empirical studies demonstrated that the exponent of the Arrhenius's power-law
324 formula falls within the range of 0.1–0.5 (Lomolino, 2001), which gives a slightly asymptotic
325 character to the fitted curve. Practically, it means that drastic increase in species numbers
326 cannot be expected with increasing habitat size. Our findings are in line with this
327 phenomenon, because despite cumulative areas of SS lakes were smaller than that of the
328 single large ones, richness of SS lakes was higher than that of SL lakes. However, one
329 exception occurred both in case of phytoplankton and benthic diatoms. This can be partly
330 explained by the above mentioned methodological limitations, but other explanations should
331 also be considered. Using a large dataset, Várбірó et al. (2017) demonstrated that the shape of
332 the SAR for phytoplankton is hump shaped, having a maximum in richness about 10^5 - 10^6 m²
333 range. Water bodies at this size range are exposed to moderate wind action and have an

334 extensive macrophyte belt; conditions which help the development of various microhabitats
335 for the phytoplankters. In large lakes, the wind induced turbulences homogenize the water
336 both horizontally and vertically creating a quasi uniform aquatic habitat. This phenomenon
337 was called the Large Lake Effect (LLE), and this seems to explain our findings that the lowest
338 values appeared in the largest size category.

339 Although dispersion ability of benthic taxa is lower than that of the planktic ones (Wetzel et
340 al., 2012), comparing to those groups where because of the obligate sexual reproduction mate
341 limitation exists (Havel and Shurin, 2004) both groups of microalgae are very good dispersers
342 (Padisák et al., 2016). Therefore, dispersal limitation is not a crucial factor affecting diversity
343 in microalgal meta-communities, instead, environmental filtering and demographic
344 stochasticity are those processes that determine the fate of colonizers in the habitats (Leibold
345 and Chase, 2017). Theoretically, the large area would benefit the colonization of habitats, but
346 size is a relative “notion” for algae, and very small habitats can satisfy the spatial needs of
347 various groups (Borics et al., 2016). The fact that ESR_{SS} was higher than ESR_{SL} clearly
348 highlighted that the species pool of the SS lakes cannot be considered as a subset of the SL
349 lake. Based on the logic proposed by Baselga (2010), in these situations the high species
350 turnover and the local heterogeneities maintain the compositional differences among the small
351 habitats, and contribute to the larger cumulative species and functional richness both in case
352 of phytoplankton and benthic diatoms.

353 The large within group diversity of the phytoplankton and the benthic diatoms, and the good
354 dispersal capabilities of taxa might occasionally result in species rich, but functionally
355 redundant assemblages. Therefore it is necessary to interpret the background of the SLOSS
356 dilemma at functional level. Functional richness can be a useful measure of ecosystem
357 complexity, which is determined by system attributes like amount of available resources,
358 isolation, habitat size, position of the system on the successional sequence, or random

359 processes e.g. colonization history and disturbances (Persson et al., 1996; Kitching, 2001;
360 Post 2002). These attributes has pronounced influence on the food-chain length, which in this
361 case can be considered as a top-down effect on the primary producers. Several field and
362 laboratory studies demonstrated that both planktic and benthic grazers prefer certain group of
363 algae (Parsons et al., 1967; Pimm and Kitching, 1987; Gresens and Lowe, 1994; Sommer,
364 1999; Kagami et al., 2002), and this preferential grazing contributes to maintain higher
365 complexity. Although an increasing complexity of water bodies could be demonstrated along
366 the size gradient (Fig. 8 A and Fig. 8 B), the functional composition of both algal groups
367 indicates, that this increasing complexity exists at the level of the whole size range ($10^{-2} - 10^7$
368 m^2). The results supported our third hypothesis, however, differences in habitat complexity
369 (number of FGs) between the adjacent size groups were not considerable, especially in the
370 case of benthic algal assemblages. An exception to this rule was the $10^2 - 10^3 m^2$ size range,
371 where considerably higher FG richness was found in $10^3 m^2$ water bodies than in the smaller
372 ones both for benthic diatoms and phytoplankters. Typically, planktic diatoms were missing
373 from the bomb crater ponds and from the small pools, resulting in a slightly decreasing
374 complexity here. In contrast, FGs tolerating the drying up of waters (e.g. motile diatoms, or
375 codon T) (Holzinger et al., 2010; Lukács et al., 2018; B-Béres et al., 2019), were
376 characteristics in these small sized ponds and pools. The fact however, that the number of FGs
377 was almost equal in the adjacent size categories (both in the case of phytoplankton and
378 benthic diatoms) strongly implies that higher ESR_{SS} values can be explained by the non-
379 nested nature of the species pool in the smaller water bodies, that is, identical FGs were
380 represented by different species in these waters.

381 The SLOSS debate inevitably attracted many theoretical approaches and explanations, and the
382 roots of this dilemma are deeply embedded in conservation management and landscape
383 planning. Although a popular view is, that protection of larger sized areas is better

384 (Tscharntke et al., 2002) investigations of different sized habitats and different animal and
385 plant groups revealed that there are arguments on “both sides of the SLOSS-debate”
386 (Tscharntke et al., 2002; Moussaoui and Auger, 2015). There is no doubt, fragmented
387 landscape is a common phenomenon worldwide, and creation of large, contiguous protected
388 areas is only rarely feasible (Gaz and Garcia-Boyero, 1996). However, as it was shown by a
389 number of studies (Tscharntke et al., 2002; Hokkanen et al., 2009; Rösch et al., 2015), in
390 certain cases, small habitats can be as valuable as larger sized areas. It is especially true for
391 small bodied organisms such as insects, snails or birds (Tscharntke et al., 2002). The results of
392 our study are not only in line with these previous findings, but demonstrate that for two
393 important microscopic aquatic groups, the higher conservational value of SS water bodies is
394 valid through the whole range of the area gradient. It is evitable, that from a practical point of
395 view, the conservation relevance of the water bodies of less than a few square meters is
396 negligible, thus, in respect to the 10^{-2} - 10^0 m² size range, our results could be considered
397 theoretical curiosities. However, in Hungary, after the large river regulations of the 19th
398 century, the formerly extended bogs and marshlands disappeared almost entirely, and the
399 biota of these ecosystems now survives in the remaining small bog-pools, that mostly are not
400 larger than 10^2 - 10^3 m² (Borics et al., 1998, 2003). While the Water Framework Directive
401 (2000) requires the achievement of good ecological status for all natural standing water bodies
402 larger than 50 hectares in Europe, smaller aquatic habitats do not belong under the umbrella
403 of this legislative approach. Therefore those small water bodies that are not parts of Natura
404 2000 sites are especially threatened, and need special consideration.

405

406 **5. Conclusions**

407 Results of the present study supported the view that microalgal species richness of several
408 small water bodies exceeds that of a single large one. These results are valid almost for the
409 entire scale of the area gradient, and for both phytoplankton and benthic diatoms.

410 Practical importance of these results is, that it draws attention to the fact that from a nature
411 conservation point of view, water bodies with very small areas might have relevant
412 conservational values.

413

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419

420 **7. Author contributions**

421 ÁB wrote the manuscript. GV and EÁKK carried out the statistical analyses. VBB, ÉÁKK
422 and KTK provided data. GB raised the topic, and helped the first author during the whole
423 course of research and writing of the manuscript. All authors gave final approval for
424 publication.

425

426 **8. References**

427 Arrhenius, O., 1921. Species and area. *J. Ecol.* 9, 95–99. [https://doi.org/ 10.2307/2255763](https://doi.org/10.2307/2255763).

428 Astorga, A., Death, R., Death, F., Paavola, R., Chakraborty, M., Muotka, T., 2014. Habitat
429 heterogeneity drives the geographical distribution of beta diversity: the case of New
430 Zealand stream invertebrates. *Ecol. Evol.* 4, 2693– 2702. <https://doi.org/10.1002/ece3.1124>

431 Báldi, A., 2008. Habitat heterogeneity overrides the species–area relationship. *J. Biogeogr.*
432 35, 675–681. <https://doi.org/10.1111/j.1365-2699.2007.01825.x>

433 Baselga, A., 2010. Partitioning the turnover and nestedness components of beta diversity.
434 *Global. Ecol. Biogeogr.* 19, 134-143. <https://doi.org/10.1111/j.1466-8238.2009.00490.x>

435 B-Béres, V., Lukács, Á., Török, P., Kókai, Zs., Novák, Z., T-Krasznai, E., Tóthmérész, B.,
436 Bácsi, I. 2016. Combined eco-morphological functional groups are reliable indicators of
437 colonisation processes of benthic diatom assemblages in a lowland stream. *Ecol. Ind.* 64,
438 31–38. <https://doi.org/10.1016/j.ecolind.2015.12.031>

439 B-Béres V., Török, P., Kókai, Zs., Lukács, Á., T-Krasznai, E., Tóthmérész, B., Bácsi, I.,
440 2017. Ecological background of diatom functional groups: Comparability of classification
441 systems. *Ecol. Ind.* 82, 183–188. <https://doi.org/10.1016/j.ecolind.2017.07.007>

442 B-Béres, V., Tóthmérész, B., Bácsi, I., Borics, G., Abonyi, A., Tapolczai, K., Rimet, F.,
443 Bouchez, A., Várbíró, G., Török, P., 2019. Autumn drought drives functional diversity of
444 benthic diatom assemblages of continental streams. *Adv. Water Resour.* 126, 129–136.
445 <https://doi.org/10.1016/j.advwatres.2019.02.010>

446 Berthon, V., Bouchez, A., Rimet, F., 2011. Using diatom life–forms and ecological guilds to
447 assess organic pollution and trophic level in rivers: a case study of rivers in south–eastern
448 France. *Hydrobiologia* 673, 259–271. <https://doi.org/10.1007/s10750-011-0786-1>

449 Bolgovics, Á., Ács, É., Várbíró, G., Görgényi, J., Borics, G., 2016. Species area relationship
450 (SAR) for benthic diatoms: a study on aquatic islands. *Hydrobiologia.* 764, 91-102.
451 <https://doi.org/10.1007/s10750-015-2278-1>

452 Borics, G., Padisák, J., Grigorszky, I., Oldal, I., Péterfi, L.S., Momeu, L., 1998. Green algal
453 flora of the acidic bog-lake, Balata-to, SW Hungary. *Biologia.* 53, 457-465.

454 Borics, G., Tóthmérész, B., Grigorszky, I., Padisák, J., Várbíró, G., Szabó, S., 2003. Algal
455 assemblage types of bog-lakes in Hungary and their relation to water chemistry,

456 hydrological conditions and habitat diversity. In *Phytoplankton and Equilibrium Concept:*
457 *The Ecology of Steady-State Assemblages.* Springer, Dordrecht. p:145-155.

458 Borics, G., Tóthmérész, B., Várbíró, G., Grigorszky, I., Czébely, A., Görgényi, J., 2016.
459 Functional phytoplankton distribution in hypertrophic systems across water body size.
460 *Hydrobiologia.* 764, 81-90. <https://doi.org/10.1007/s10750-015-2268-3>

461 Borics, G., Várbíró, G., Grigorszky, I., Krasznai, E., Szabó, S., Kiss, K.T., 2007. A new
462 evaluation technique of potamo-plankton for the assessment of the ecological status of
463 rivers. *Arch. Hydrobiol. Suppl.* 17, 465–486. <https://doi.org/10.1127/lr/17/2007/465>

464 Buckland, S.T., Studeny, A.C., Magurran, A.E. and Newson, S.E., 2011. Biodiversity
465 monitoring: the relevance of detectability. in: A Magurran, A., McGill, B., (Eds.),
466 *Biological Diversity: Frontiers in Measurement and Assessment.* Oxford University Press,
467 pp. 25-36.

468 Chao, A., Gotelli, N.J., Hsieh, T.C., Sander, E.L., Ma, K.H., Colwell, R.K., Ellison, A.M.,
469 2014. Rarefaction and extrapolation with Hill numbers: a framework for sampling and
470 estimation in species diversity studies. *Ecol. Monogr.* 84, 45-67.
471 <https://doi.org/10.1890/13-0133.1>

472 Connor, E.F., McCoy, E., 1979. The statistics and biology of the species-area relationship.
473 *Am. Nat.* 113, 791-833. <http://www.jstor.org/stable/2460305>.

474 Diamond, J.M., 1975. The island dilemma: lessons of modern biogeographic studies for the
475 design of natural reserves. *Biol. Conserv.* 7, 129-146. [https://doi.org/10.1016/0006-](https://doi.org/10.1016/0006-3207(75)90052-X)
476 [3207\(75\)90052-X](https://doi.org/10.1016/0006-3207(75)90052-X)

477 Dodson, S.I., 1992. Predicting Crustacean zooplankton species richness. *Limnol. Oceanogr.*
478 37, 848–856. <https://doi.org/10.4319/lo.1992.37.4.0848>

479 EC (2000) Directive 2000/60/EC of the European Parliament and of the Council of 23rd
480 October 2000 establishing a framework for Community action in the field of water policy.

481 Official Journal of the European Communities, 22 December, L 327/1. European
482 Commission, Brussels.

483 Fahrig, L., 2002. Effect of habitat fragmentation on the extinction threshold: a synthesis. *Ecol.*
484 *Appl.* 12, 346-353. [https://doi.org/10.1890/1051-0761\(2002\)012\[0346:EOHFOT\]2.0.CO;2](https://doi.org/10.1890/1051-0761(2002)012[0346:EOHFOT]2.0.CO;2)

485 Fahrig, L., 2003. Effects of habitat fragmentation on biodiversity. *Annu. Rev. Ecol. Evol. S.*
486 34, 487-515. <https://doi.org/10.1146/annurev.ecolsys.34.011802.132419>

487 Fischer, J., Lindenmayer, D.B., 2007. Landscape modification and habitat fragmentation: a
488 synthesis. *Global. Ecol. Biogeogr.* 16, 265–280. [https://doi.org/10.1111/j.1466-](https://doi.org/10.1111/j.1466-8238.2007.00287.x)
489 [8238.2007.00287.x](https://doi.org/10.1111/j.1466-8238.2007.00287.x)

490 Foley, J.A., DeFries, R., Asner, G.P., Barford, C., Bonan, G., Carpenter, S.R., Chapin, F.S.,
491 Coe, M.T., Daily, G.C., Gibbs, H.K., Helkowski, J.H., Holloway, T., Howard, E.A.,
492 Kucharik, C.J., Monfreda, C., Patz, J.A., Prentice, I.C., Ramankutty, N., Snyder, P.K..
493 2005. Global consequences of landuse. *Science.* 309, 570-574.
494 <https://doi.org/10.1126/science.1111772>

495 Gao, D., Perry, G., 2016. Detecting the small island effect and nestedness of herpetofauna of
496 the West Indies. *Ecol. Evol.* 15, 5390– 5403. <https://doi.org/10.1002/ece3.2289>

497 Gaz, A., Garcia-Boyer, A., 1996. The SLOSS-dilemma: a butterfly case study. *Biodivers.*
498 *Conserv.* 5, 493-502. <https://doi.org/10.1007/BF00056393>

499 Gibb, H., Hochuli, D.F., 2002. Habitat fragmentation in an urban environment: large and
500 small fragments support different arthropod assemblages. *Biol. Conserv.* 106, 91–100.
501 [https://doi.org/10.1016/S0006-3207\(01\)00232-4](https://doi.org/10.1016/S0006-3207(01)00232-4)

502 Gilpin, M.E., Hanski, I.A., 1991. Metapopulation dynamics: brief history and conceptual
503 domain. *Biol. J. Linn. Soc.* 42, 3–16. <https://doi.org/10.1111/j.1095-8312.1991.tb00548.x>

504 Gresens, S.E. Lowe, R.L., 1994. Periphyton patch preference in grazing chironomid larvae. *J.*
505 *N. Am. Benthol. Soc.*, 13, 89–99. <https://doi.org/10.2307/1467269>

506 Görgényi, J., Tóthmérész, B., Várbiro, G., Abonyi, A., T-Krasznai, E., B-Béres V., . Borics,
507 G., 2019. Contribution of phytoplankton functional groups to the diversity of a eutrophic
508 oxbow lake Hydrobiologia ACCEPTED in Hydrobiologia. <https://doi.org/10.1007/s10750->
509 018-3878-3

510 Havel, J.E., Shurin, J.B., 2004. Mechanisms, effects, and scales of dispersal in freshwater
511 zooplankton. *Limnol Oceanogr.* 49, 1229-1238.
512 https://doi.org/10.4319/lo.2004.49.4_part_2.1229

513 Harrison, S., Ross, S.J., Lawton, J.H., 1992. Beta-diversity on geographic gradients in Britain.
514 *J. Anim. Ecol.* 61, 151–158. <https://doi.org/10.2307/5518>

515 Hokkanen, P.J., Kouki, J., Komonen, J., 2009. Nestedness. SLOSS and conservation networks
516 of boreal herb-rich forests. *Appl. Veg. Sci.* 12, 295–303. <https://doi.org/10.1111/j.1654->
517 [109X.2009.01031.x](https://doi.org/10.1111/j.1654-109X.2009.01031.x)

518 Holzinger, A., Tschaikner, A., Remias, D., 2010. Cytoarchitecture of the desiccation-tolerant
519 green alga *Zygonium ericetorum*. *Protoplasma* 243, 15–24. DOI 10.1007/s00709-009-
520 0048-5

521 Honnay, O., Hermy, M., Coppin, P., 1999. Effects of area, age and diversity of forest patches
522 in Belgium on plant species richness, and implications for conservation and reforestation.
523 *Biol. Conserv.* 87, 73–84. [https://doi.org/10.1016/S0006-3207\(98\)00038-X](https://doi.org/10.1016/S0006-3207(98)00038-X)

524 Hsieh, T.C., Ma, K.H., Chao, A., 2013. iNEXT online: interpolation and extrapolation
525 (Version 1.0) [Software]. Available from [http://chao.stat.nthu.edu.tw/blog/software-](http://chao.stat.nthu.edu.tw/blog/software-download)
526 [download.](http://chao.stat.nthu.edu.tw/blog/software-download)

527 Hylander, K., Nilsson, C., Gunnar-Jonsson, B., Göthner, T., 2005. Differences in habitat
528 quality explain nestedness in a land snail meta- community. *Oikos.* 108, 351-361.
529 <https://doi.org/10.1111/j.0030-1299.2005.13400.x>

530 Kagami, M., Yoshida, T., Gurung, T. and Urabe, J., 2002. Direct and indirect effects of
531 zooplankton on algal composition in in situ grazing experiments. *Oecologia*, 133(3), 356–
532 363. <https://doi.org/10.1007/s00442-002-1035-0>

533 Le Roux, D.S., Ikin, K., Lindenmayer, D.B., Manning, A.D., Gibbons, P., 2015. Single large
534 or several small? Applying biogeographic principles to tree-level conservation and
535 biodiversity offsets. *Biol. Conserv.* 191, 558–566.
536 <https://doi.org/10.1016/j.biocon.2015.08.011>

537 Leibold, M.A., Chase, J.M., 2017. *Metacommunity ecology* (Vol. 59). Princeton University
538 Press.

539 Leibold, M.A., Holyoak, M., Mouquet, N., Amarasekare, P., Chase, J.M., Hoopes, M.F., Holt,
540 R.D., Shurin, J.B., Law, R., Tilman, D., Loreau, M., Gonzalez, A., 2004. The
541 metacommunity concept: a framework for multi-scale community ecology. *Ecol. Lett.* 7,
542 601–613. <https://doi.org/10.1111/j.1461-0248.2004.00608.x>

543 Liess, A., Hillebrand, H., 2004. Invited review: direct and indirect effects in herbivore
544 periphyton interactions. *Fund. Appl. Limnol.* 159, 433–453. DOI: 10.1127/0003-
545 9136/2004/0159-0433

546 Lindenmayer, D.B., Wood, J., McBurney, L., Blair, D., Banks, S.C., 2015. Single large versus
547 several small: The SLOSS debate in the context of bird responses to a variable retention
548 logging experiment. *Forest Ecol. Manag.* 339, 1–10.
549 <https://doi.org/10.1016/j.foreco.2014.11.027>

550 Lomolino, M.V., Weiser, M.D., 2001. Towards a more general species–area relationship:
551 diversity on all islands, great and small. *J. Biogeogr.* 28, 431–445.
552 <https://doi.org/10.1046/j.1365-2699.2001.00550.x>

553 Lomolino, M.V., 2001. The species–area relationship: new challenges for an old pattern.
554 *Prog. Phys. Geog.* 25, 1–21. <https://doi.org/10.1177/030913330102500101>

555 Lukács, Á., Kókai, Zs., Török, P., Bácsi, I., Borics, G., Várbíró, G., T-Krasznai, E.,
556 Tóthmérész, B., B-Béres, V., 2018. Colonisation processes in benthic algal communities
557 are well reflected by functional groups. *Hydrobiologia* 823, 231–245.
558 <https://doi.org/10.1007/s10750-018-3711-z>

559 Matias, M.G., Underwood, A.J., Hochuli, D.F., Coleman, R.A., 2010. Independent effects of
560 patch size and structural complexity on diversity of benthic macroinvertebrates. *Ecology*.
561 91, 1908-1915. <https://doi.org/10.1890/09-1083.1>

562 Matthews, T.J., Guilhaumon, F., Triantis, K.A., Borregaard, M.K., Whittaker, R.J., 2016. On
563 the form of species–area relationships in habitat islands and true islands. *Global Ecol.*
564 *Biogeogr.* 25, 847–858. <https://doi.org/10.1111/geb.12269>

565 Moussaoui, A., Auger, P., 2015. Simple fishery and marine reserve models to study the
566 SLOSS problem. *ESAIM Proc. Surv.* 49, 78-90. <https://doi.org/10.1051/proc/201549007>

567 OECD 1982. Eutrophication of Waters. Monitoring, assessment and control. Final Report,
568 OECD cooperative programme on monitoring of inland waters (Eutrophication control),
569 Environment Directorate. – OECD, Paris, pp.154.

570 Padisák, J., Borics, G., Grigorszky, I. and Soroczki-Pintér, E., 2006. Use of phytoplankton
571 assemblages for monitoring ecological status of lakes within the Water Framework
572 Directive: the assemblage index. *Hydrobiologia*, 553(1), 1–14.

573 Padisák, J., Crossetti, L.O., Naselli-Flores, L., 2009. Use and misuse in the application of the
574 phytoplankton functional classification: a critical review with updates. *Hydrobiologia*. 621,
575 1–19. <https://doi.org/10.1007/s10750-008-9645-0>

576 Padisák, J., Vasas, G., Borics, G., 2016. Phycogeography of freshwater phytoplankton:
577 traditional knowledge and new molecular tools. *Hydrobiologia*. 764, 3–27.
578 <https://doi.org/10.1007/s10750-015-2259-4>

579 Parsons, T.R., LeBrasseur, R.J., Fulton, J.D., 1967. Some observations on the dependence of
580 zooplankton grazing on the cell size and concentration of phytoplankton blooms. *J.*
581 *Oceanogr. Soc. Japan*, 23, 10-17. DOI: [10.5928/kaiyou1942.23.10](https://doi.org/10.5928/kaiyou1942.23.10)

582 Passy, S., 2007. Diatom ecological guilds display distinct and predictable behavior along
583 nutrient and disturbance gradients in running waters. *Aquat. Bot.* 86, 171–178.
584 <https://doi.org/10.1016/j.aquabot.2006.09.018>

585 Pimm, S.L. Kitching, R.L., 1987. The determinants of food chain lengths. *Oikos*, 302–307.
586 <https://www.jstor.org/stable/3565490>

587 Reynolds, C.S., Huszár, V., Kruk, C., Naselli-Flores, L., Melo, S., 2002. Towards a functional
588 classification of the freshwater phytoplankton. *J. Plankton Res.* 24, 417–428.
589 <https://doi.org/10.1093/plankt/24.5.417>

590 Rimet, F., Bouchez, A., 2012. Life-forms, cell-sizes and ecological guilds of diatoms in
591 European rivers. *Knowl. Manag. Aquat. Ec.* 406, 01.
592 <https://doi.org/10.1051/kmae/2012018>

593 Rösch, V., Tschardtke, T., Scherber, C., Batáry, P., 2015. Biodiversity conservation across
594 taxa and landscapes requires many small as well as single large habitat fragments.
595 *Oecologia*. 179, 209–222. <https://doi.org/10.1007/s00442-015-3315-5>

596 RStudio. 2012. RStudio: Integrated development environment for R (Version 0.97)
597 [Computer software]. Boston, MA. Available from: <http://www.rstudio.org/>

598 Salmaso, N., Naselli-Flores, L., Padisák, J., 2015. Functional classifications and their
599 application in phytoplankton ecology. *Freshwater Biol.* 60, 603–619.
600 <https://doi.org/10.1111/fwb.12520>

601 Simberloff, D., Abele, L.G., 1976. Island biogeography theory and conservation practice.
602 *Science*. 191, 285-286. <https://doi.org/10.1126/science.191.4224.285>

603 Smith, V.H., Foster, B.L., Grover, J.P., Holt, R.D., Leibold, M.A., deNoyelles, F. Jr., 2005.
604 Phytoplankton species richness scales consistently from laboratory microcosms to the
605 world's oceans. *PNAS*. 102, 4393–4396. <https://doi.org/10.1073/pnas.0500094102>

606 Soininen, J., Jamoneau, A., Rosebery, J., Passy, S.I., 2016. Global patterns and drivers of
607 species and trait composition in diatoms. *Global Ecol. Biogeogr.* 8, 940-950.
608 <https://doi.org/10.1111/geb.12452>

609 Soininen, J., Meier, S., 2014. Phytoplankton richness is related to nutrient availability, not to
610 pool size, in a subarctic rock pool system. *Hydrobiologia* 740, 137–145.
611 <https://doi.org/10.1007/s10750-014-1949-7>

612 Sommer, U., 1999. The susceptibility of benthic microalgae to periwinkle (*Littorina littorea*,
613 Gastropoda) grazing in laboratory experiments. *Aquatic botany*, 63(1), 11–21.
614 [https://doi.org/10.1016/S0304-3770\(98\)00108-9](https://doi.org/10.1016/S0304-3770(98)00108-9)

615 Stenger-Kovács, Cs., Körmendi, K., Lengyel, E., Abonyi, A., Hajnal, É., Szabó, B., Buczkó,
616 K., Padisák, J., 2018. Expanding the trait-based concept of benthic diatoms: Development
617 of trait- and species-based indices for conductivity as the master variable of ecological
618 status in continental saline lakes. *Ecol. Ind.* 95, 63-74.
619 <https://doi.org/10.1016/j.ecolind.2018.07.026>

620 Striebel, M., Singer, G., Stibor, H. and Andersen, T., 2012. “Trophic overyielding”:
621 Phytoplankton diversity promotes zooplankton productivity. *Ecology*, 93(12), 2719-2727.

622 Szabó, B., Lengyel, E., Padisák, J., Stenger-Kovács, Cs., 2018. Benthic diatom
623 metacommunity across small freshwater lakes: driving mechanisms, β -diversity and
624 ecological uniqueness. *Hydrobiologia* 828, 183-198. [https://doi.org/10.1007/s10750-018-](https://doi.org/10.1007/s10750-018-3811-9)
625 [3811-9](https://doi.org/10.1007/s10750-018-3811-9)

626 Tapolczai, K., Bouches, A., Stenger-Kovács, Cs., Padisák, J., Rimet, F., 2016. Trait-based
627 ecological classifications for benthic algae: review and perspectives. *Hydrobiologia* 776 ,
628 1-17. <https://doi.org/10.1007/s10750-016-2736-4>

629 Tjørve, E., 2010. How to Resolve the SLOSS debate: Lessons from Species-diversity Models.
630 *J. Theor. Biol.* 264, 604-612. <https://doi.org/10.1016/j.jtbi.2010.02.009>

631 Tjørve, E., 2003. Shapes and functions of species–area curves: a review of possible models. *J.*
632 *Biogeogr.* 30, 827–835. <https://doi.org/10.1046/j.1365-2699.2003.00877.x>

633 Török, P., T- Krasznai, E., B- Béres, V., Bácsi, I., Borics, G., Tóthmérész, B., 2016.
634 Functional diversity supports the biomass–diversity humped- back relationship in
635 phytoplankton assemblages. *Funct. Ecol* 30, 1593-1602. [https://doi.org/10.1111/1365-](https://doi.org/10.1111/1365-2435.12631)
636 [2435.12631](https://doi.org/10.1111/1365-2435.12631)

637 Townsend, S.A., Gell, P.A., 2005. The role of substrate type on benthic diatom assemblages
638 in the Daly and Roper Rivers of the Australian wet/dry tropics. *Hydrobiologia* 548, 101-
639 115. <https://doi.org/10.1007/s10750-005-0828-7>

640 Triantis, K.A., Sfenthourakis, S., 2011. Island biogeography is not a single variable discipline:
641 the small island effect debate. *Divers Distrib.* 18, 92–96. [https://doi.org/10.1111/j.1472-](https://doi.org/10.1111/j.1472-4642.2011.00812.x)
642 [4642.2011.00812.x](https://doi.org/10.1111/j.1472-4642.2011.00812.x)

643 Tscharnkte, T., Steffan-Dewenter, I., Kruess, A., Thies, C., 2002. Contribution of small
644 habitat fragments to conservation of insect communities of grassland-cropland landscapes.
645 *Ecol. Appl.* 12, 354-363. [https://doi.org/10.1890/1051-](https://doi.org/10.1890/1051-0761(2002)012[0354:COSHFT]2.0.CO;2)
646 [0761\(2002\)012\[0354:COSHFT\]2.0.CO;2](https://doi.org/10.1890/1051-0761(2002)012[0354:COSHFT]2.0.CO;2)

647 Turner, I.M., Corlett, R.T., 1996. The conservation value of small, isolated fragments of
648 lowland tropical rainforest. *Trends Ecol. Evol.* 11, 330-333. [https://doi.org/10.1016/0169-](https://doi.org/10.1016/0169-5347(96)10046-X)
649 [5347\(96\)10046-X](https://doi.org/10.1016/0169-5347(96)10046-X)

650 Utermöhl, H., 1958. Zur Vervollkommnung der quantitative Phytoplankton-Methodik. *Mitt.*
651 *Int. Ver. Limn.* 9, 1–38.

652 Várbíró, G., Görgényi, J., Tóthmérész, B., Padisák, J., Hajnal, É., Borics G., 2017. Functional
653 redundancy modifies species–area relationship for freshwater phytoplankton. *Ecol. Evol.* 7,
654 9905–9913. <https://doi.org/10.1002/ece3.3512>

655 Wetzel, R.G., de C. Bicudo, D., Ector, L., Lobo, E.A., Soininen, J., Landeiro, V.L., Bini,
656 L.M., 2012. Distance Decay of Similarity in Neotropical Diatom Communities. PlosOne 7,
657 e45071. <https://doi.org/10.1371/journal.pone.0045071>

658 Wickham, H., 2011. ggplot2. Wiley Interdisciplinary Reviews: Computational Statistics. 3,
659 180-185.

660 Wiens, J.A., 1974. Habitat heterogeneity and avian community structure in North American
661 grasslands. Am. Midl. Nat. 91, 195-213. <http://www.jstor.org/stable/2424522>.

662 Wilson, E.O., Willis, E.O., 1975. Applied biogeography. In: Ecology and Evolution of
663 Communities. p. 523-534. Cambridge, MA:Belknap Press of Harvard University.

664 [database 1] <http://www.vizugy.hu/index.php?module=vizstrat&programelemid=149>

665

666 **9. Captions**

667

668 Table 1 Sample numbers (original and estimated) considered in a given sample site. Black
 669 arrows indicate how we divided the species richness of SL lakes with species richness
 670 of SS lakes in the one smaller size category.

Size category (m ²)	The name of the water bodies	Observed number of samples in case of benthic diatoms	Observed number of samples in case of phytoplankton	Number of samples considered for the analyses (SS)	Estimated sample number (SL)
0.01 (10 ⁻²)	Bomb crater	5	5	5	
0.1 (10 ⁻¹)	Bomb crater	5	5	5	5
1 (10 ⁰)	Bomb crater	5	5	5	5
10 (10 ¹)	Bomb crater	5	5	5	5
100 (10 ²)	Bomb crater	5	5	5	5
1000 (10 ³)	Felső Darab Tisza	9	9		
	Egyekpusztakócsi mocsár (Hagymás)	5			
	Sáros-ér		3		
	Morotvaközi H-Meder, Egyek	5	5	15	
10000 (10 ⁴)	Egyeki H-Tisza, Egyek	17	11		15
	Tiszadobi Holt-Tisza, Darab Tisza	10	10	30	15
	Egyek-Kócsi Tározó, Góré	4	4		15
100000 (10 ⁵)	Tiszadobi Holt-Tisza, Falu-Tisza	15	15		30
	Tiszadobi Holt-Tisza, Malom-Tisza	34	34	45	30
	Tiszadobi Holt-Tisza, Szűcs-Tisza	15	15		30
1000000 (10 ⁶)	H-Szamos, Tunyogmatolcs+ Géberjén	23	23	60	45
10000000 (10 ⁷)	Velencei-tó	27	27		60

671

672 Table A.1 Limnological characteristics of studied lakes. Characterisation of lakes' trophic
 673 level was based on the OECD proposal (1982) (hypertrophic: TP > 100 mg/l).

674

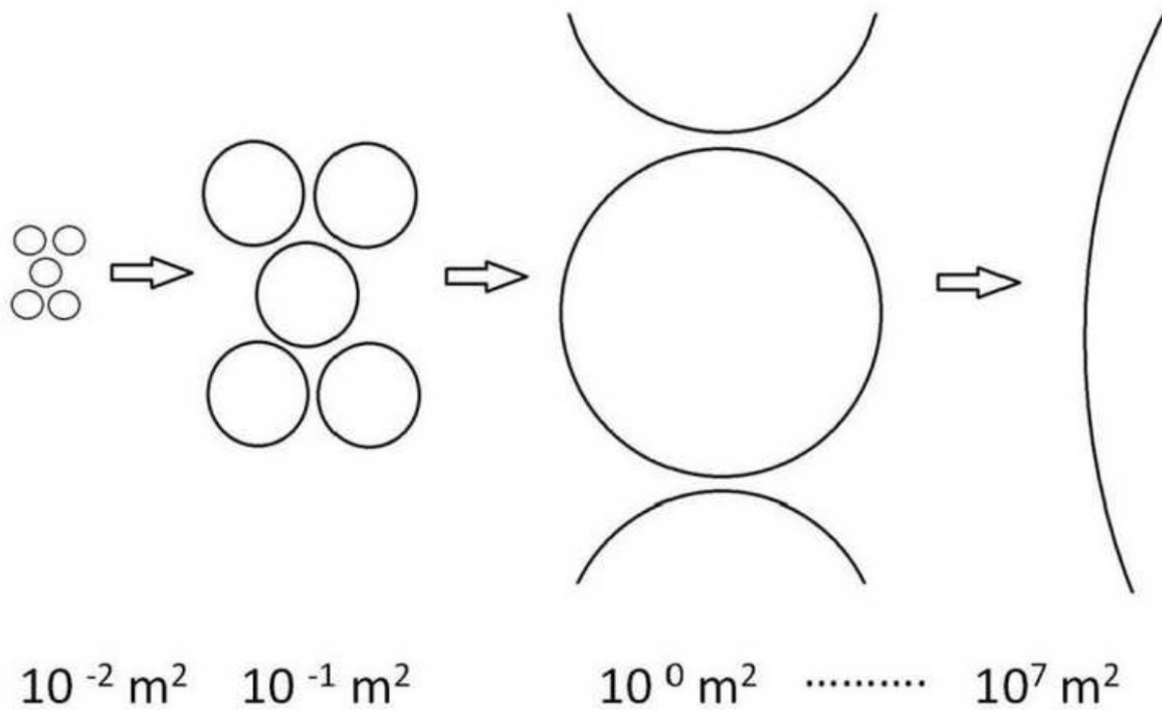
675 Table A.2 Codes of the combined eco-morphological functional groups of diatoms

676

677 Table A.3 Characteristics of the observed phytoplankton functional groups

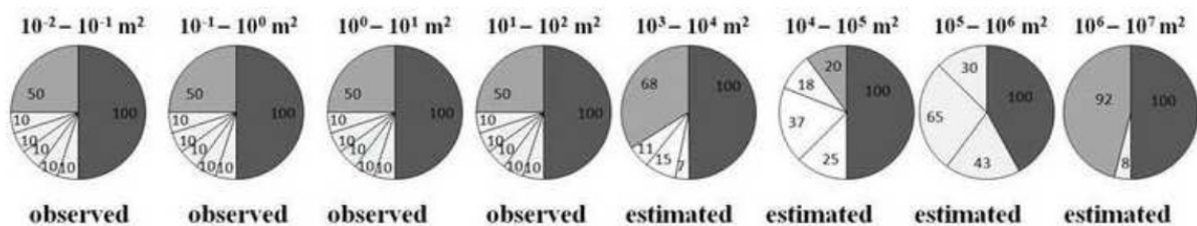
678

679 Fig. 1 Illustration of the applied study design. Circles represent the area of the water bodies.



680

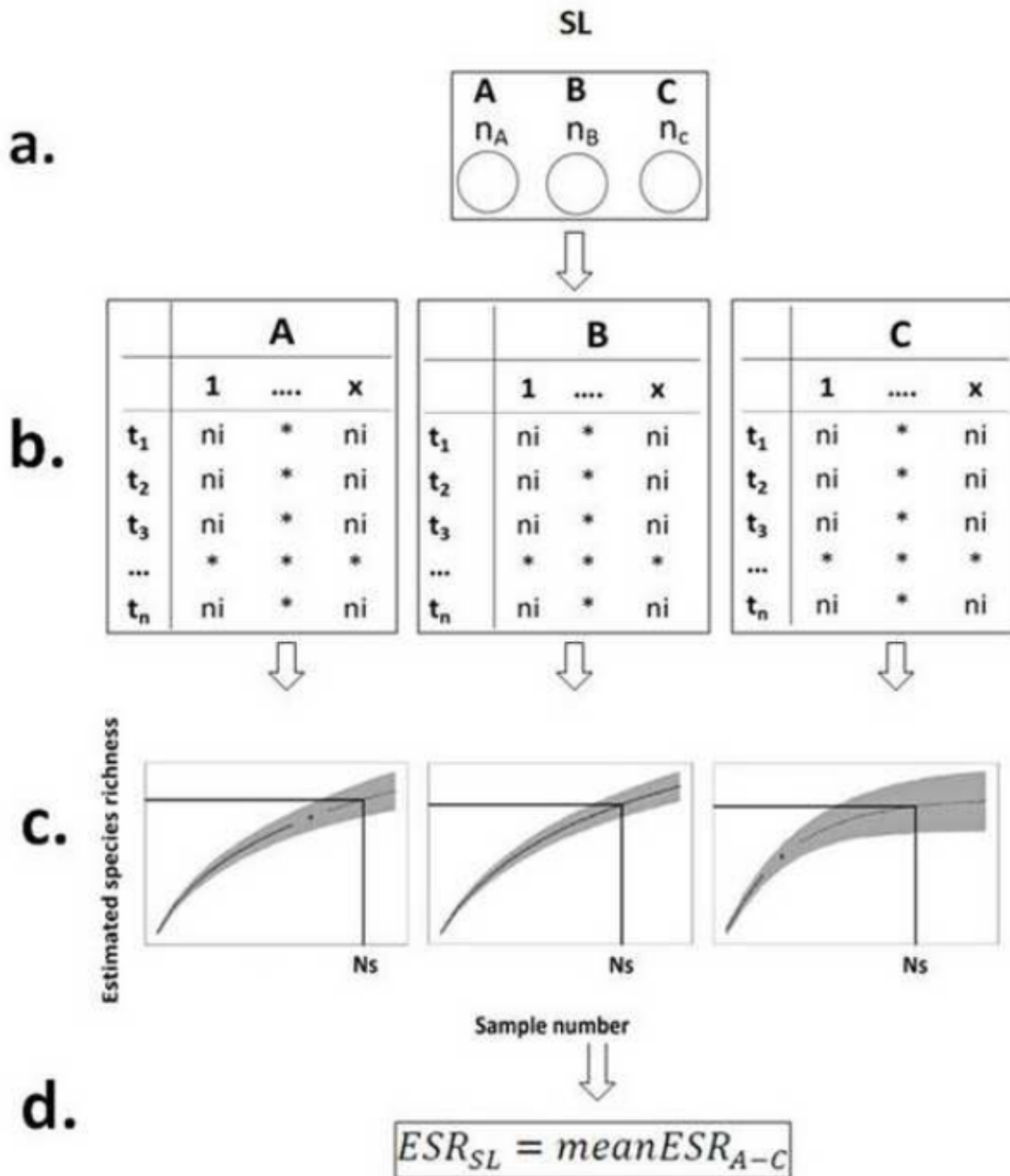
681 Fig. 2 Area covered by the SS lakes comparing to that of SL lakes. The dark grey part of the
 682 pie charts with 100 % represents the size of the SL lakes. The white parts of the pie charts
 683 show the size of the SS lakes expressed as the percentage of the area of the SL lake. Area of
 684 the SL lake was considered as the mean area of the lakes in the given size category. Numbers
 685 in the pie charts indicate the percentages covered by the small lakes. The light grey parts show
 686 the ratio of uncovered area.



687

688 Fig. 3 Calculation of the species richness for the single large (SL) lakes (SL: 10^3-10^7 m^2)
 689 within a given size category. Abbreviations: A, B, C – water bodies; n (A, B, C) – sample
 690 number; t – taxa; ni – number of individuals; ESR – estimated species richness; Ns – number

691 of samples considered during richness estimations; SL – single large; SS – several small
 692 lakes.



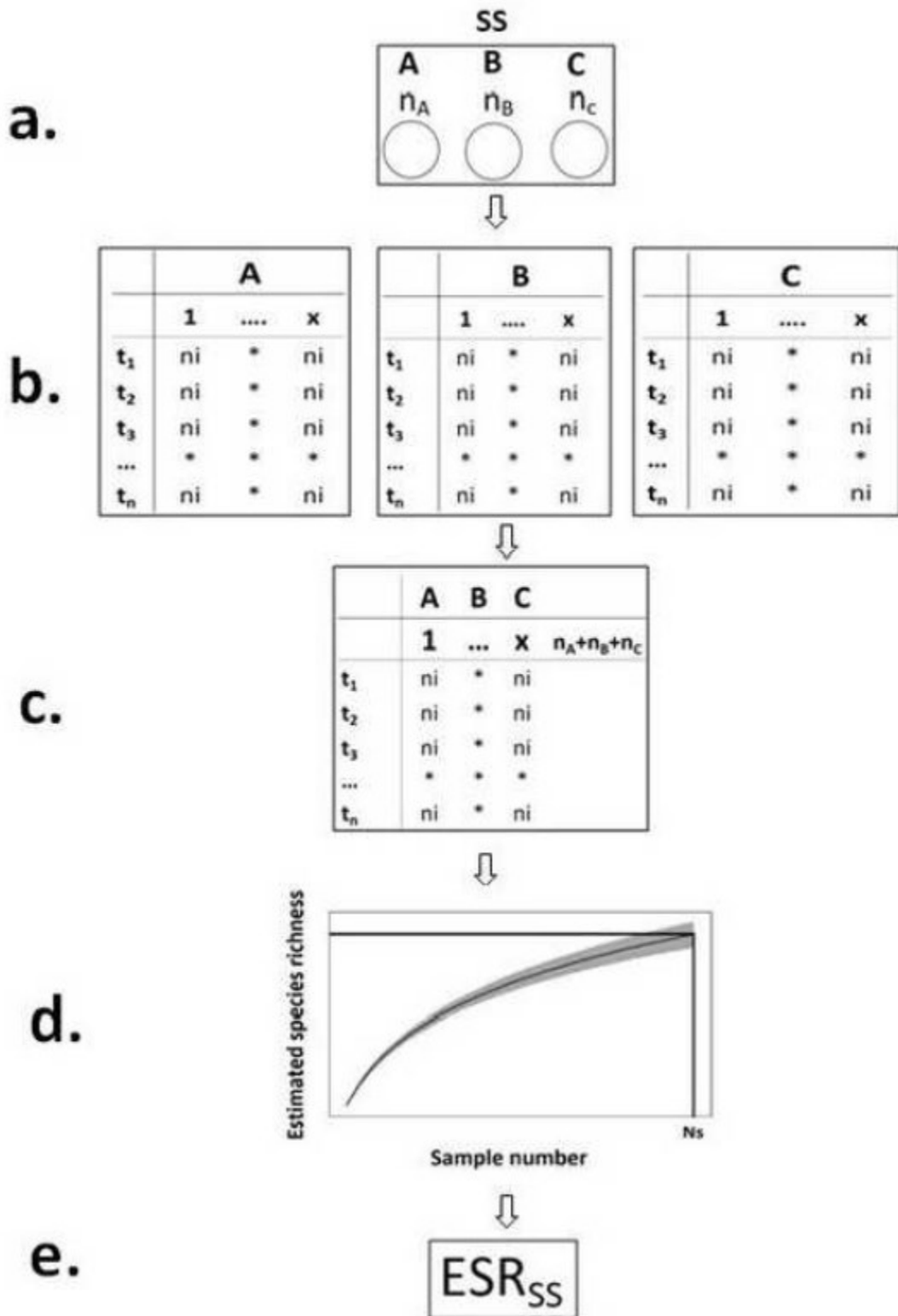
693

694 Fig. 4 Calculation of the species richness for the several small (SS) lakes (SL: 10^3 - 10^7 m²)

695 within a given size category. Abbreviations: A, B, C – water bodies; n (A, B, C) – sample

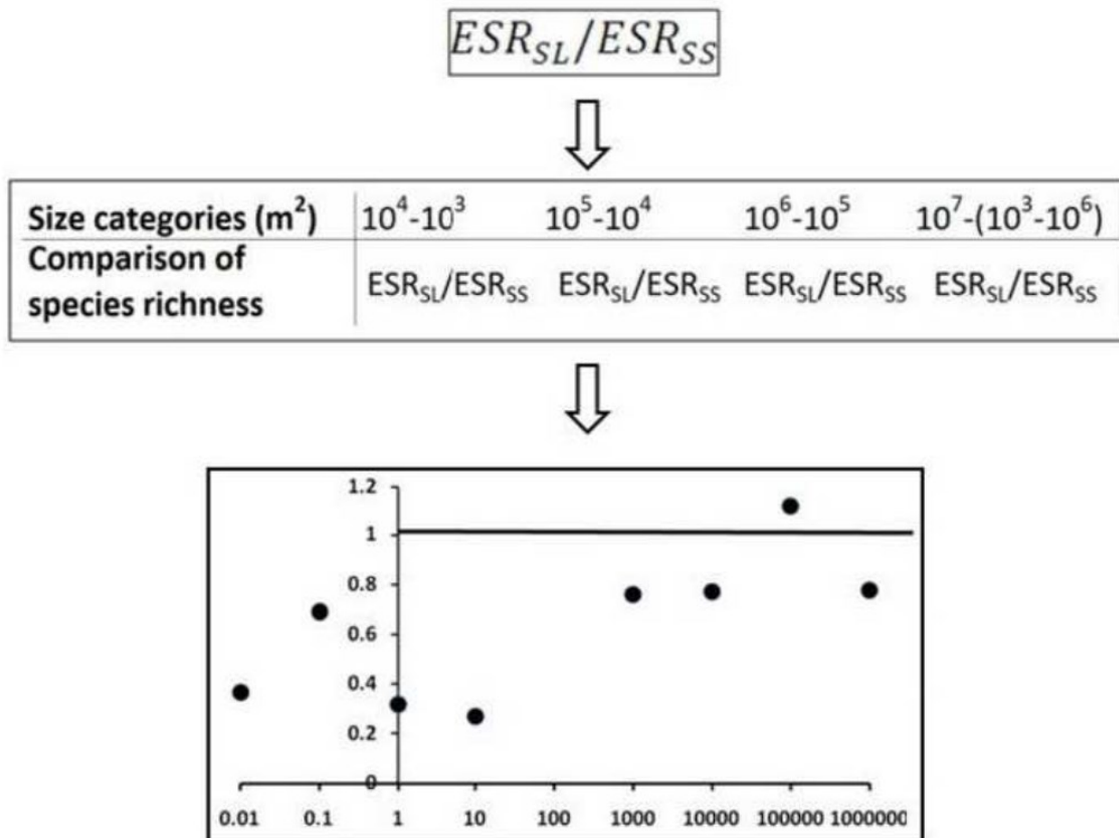
696 number; t – taxa; n_i – number of individuals; ESR – estimated species richness; N_s – number

697 of samples considered during richness estimations; SL – single large; SS – several small
 698 lakes.



699

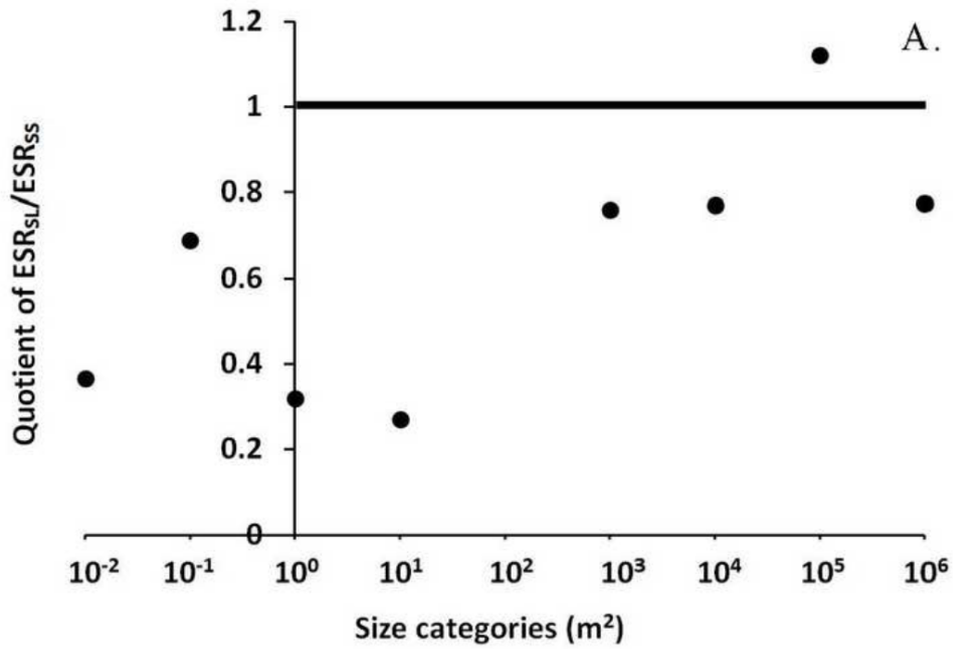
700 Fig. 5 Numerical characterisation of the SLOSS debate and its presentation in the compared
 701 water body size categories. ESR_{SL} : estimated species richness in single large lake, ESR_{SS} :
 702 estimated species richness for several small lakes.



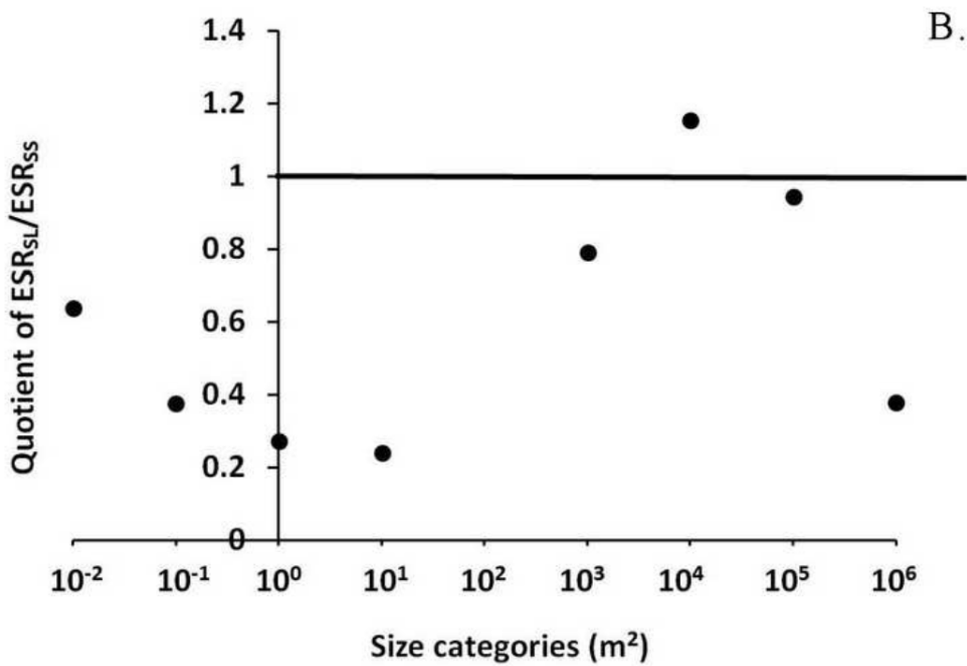
703

704

705 Fig. 6 A-B Benthic diatom and phytoplankton ESR_{SL}/ESR_{SS} values in the compared water
 706 body size categories. Values under black line show when the species richness of SS lakes
 707 were higher than in case of SL lakes, while the values above the black line mark higher
 708 species richness of SL lakes than in SS ones.



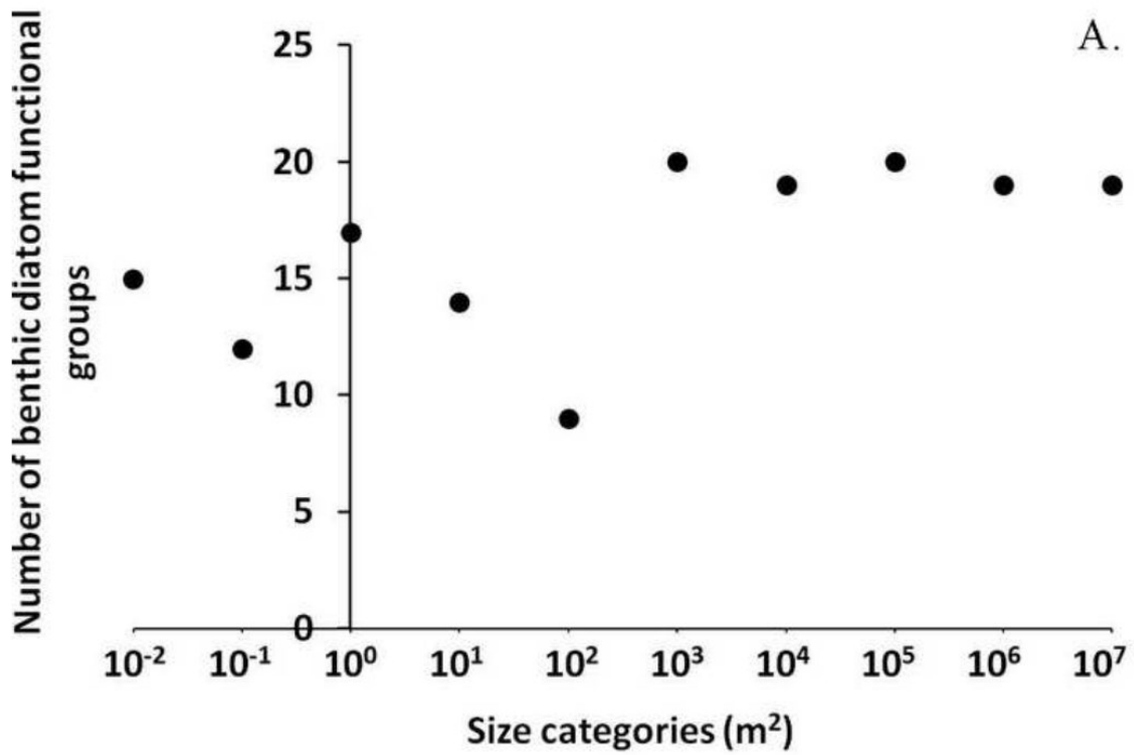
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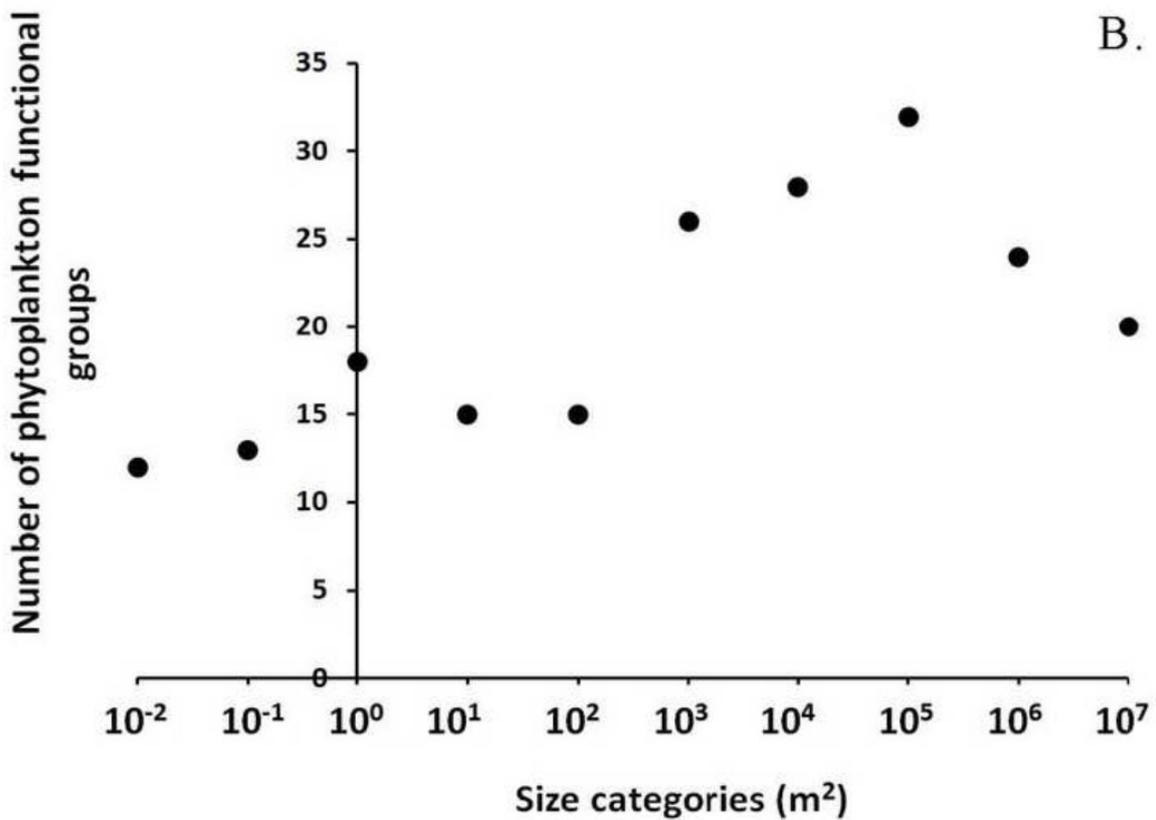
710

711 Fig. 7 A-B. Cumulative number of benthic diatom and phytoplankton FGs in the water body

712 size categories.

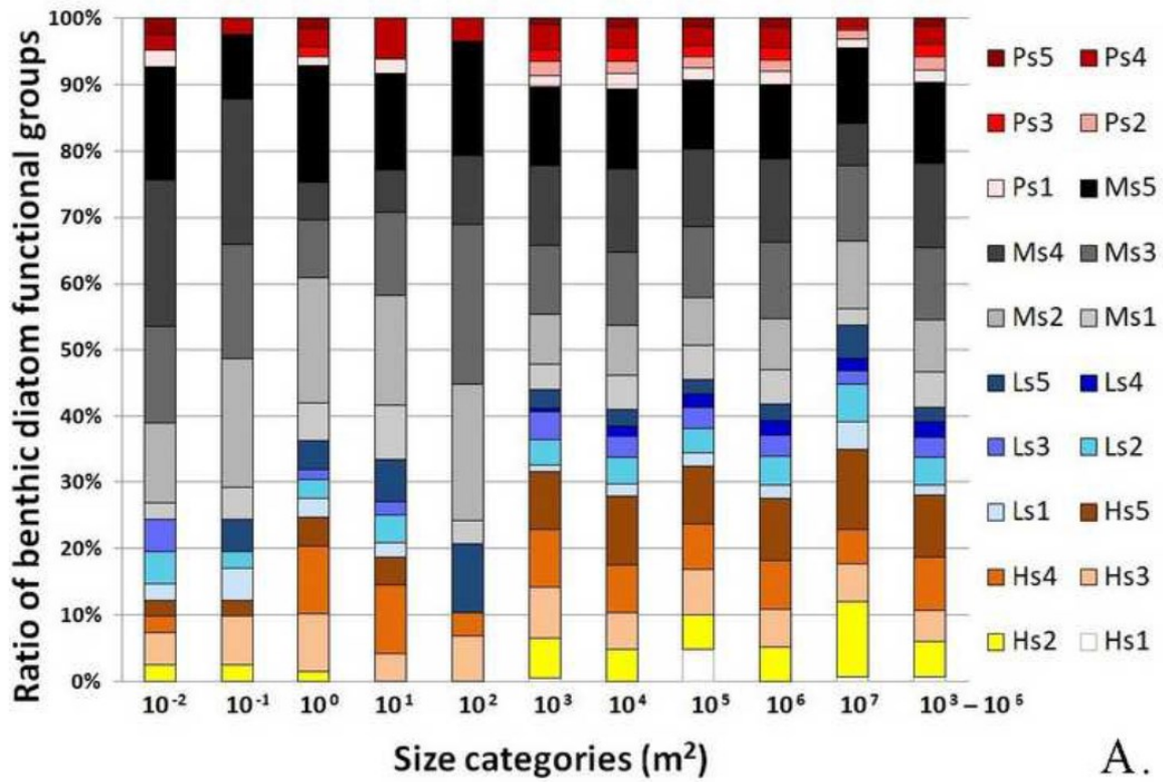


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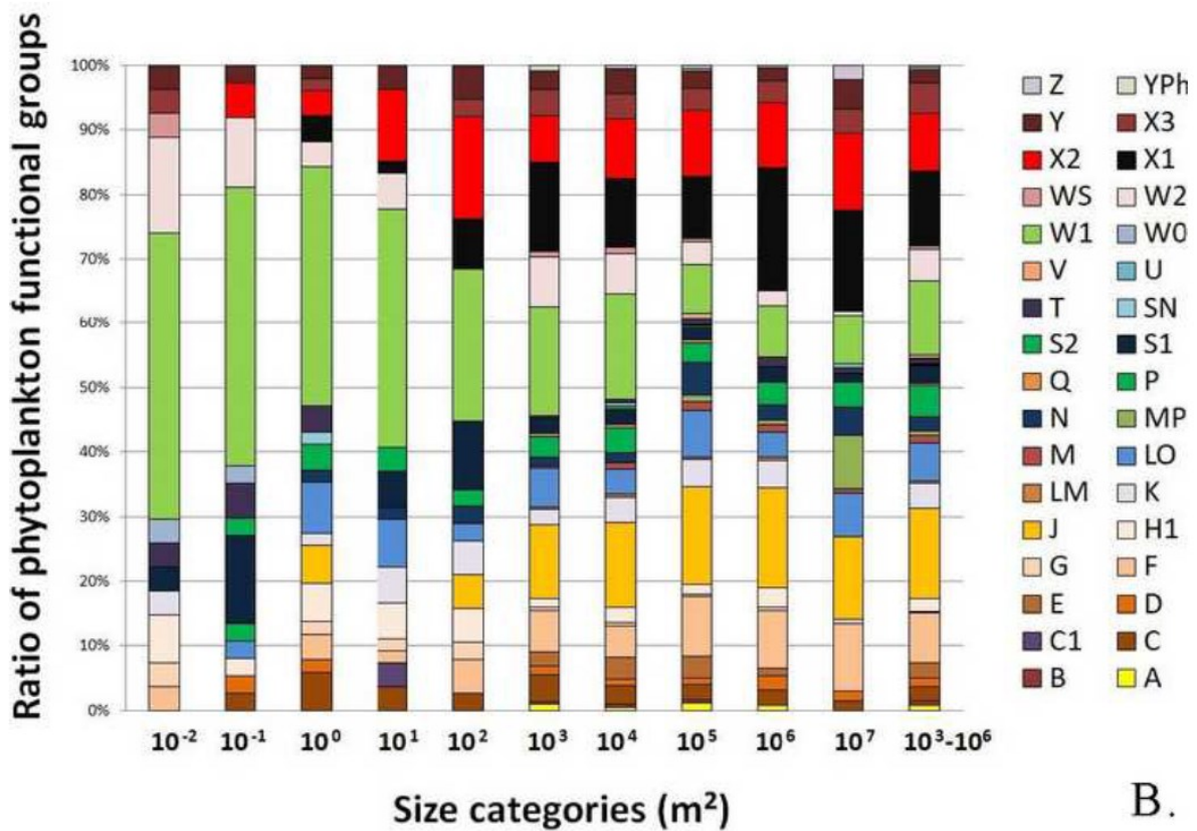


714

715 Fig. 8 A-B Relative species abundances in the functional groups of benthic diatoms and
 716 phytoplankton in the different size categories. See abbr. in Table A.1 and Table A.3



717



718