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Population divergence in aggregation and sheltering behaviour in surface and cave-adapted Asellus aquaticus (Crustacea: Isopoda)

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2	surface and cave-adapted Asellus aquaticus (Crustacea: Isopoda)
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15 Abstract

16 Aggregation (gathering together) and sheltering (hiding in cover) are basic behaviours that might reduce the risk of predation. However, both behaviours have costs, like increased 17 competition over resources and high prevalence of contact-spread parasites (aggregation) or 18 lost opportunities for foraging and mating (sheltering). Therefore, adaptive variation in these 19 behaviours is expected between populations with varying levels of predation risk. We 20 compared aggregation and sheltering in surface- (various predators) and cave-adapted (no 21 22 predator) populations of the Isopod Asellus aquaticus in a common garden experiment. Since the cave environment is constantly dark, we also tested for population variation in light-23 24 induced behavioural plasticity by applying light treatments. Variation in sheltering was explained by habitat type: cave individuals sheltered less than surface individuals. We found 25 high between-population variation in aggregation with or without shelters and their light-26 27 induced plasticity, which were unexplained by habitat type. Cave individuals habituated (decreased sheltering), while surface individuals showed sensitization (increased sheltering). 28 29 We suggest that population variation in sheltering is driven by predation, while variation in 30 aggregation must be driven by other, unaccounted environmental factors, similarly to light-31 induced behavioural plasticity. Based on habituation/sensitization patterns, we suggest that 32 predation-adapted populations are more sensitive to disturbance related to standard laboratory procedures. 33

34

35 Keywords: aggregation, Asellus aquaticus, cave adaptation, shelter use

37 Introduction

38 Behaviour is perhaps the most plastic quantitative phenotypic trait (West-Eberhard, 2003). However, despite high plasticity and the potential for a moment-by-moment optimisation to 39 the prevailing environment, geographic between-population variation within species in 40 behaviour is evident, suggesting local (genetic) adaptation in behaviour (e.g. Foster, 1999; 41 Foster & Endler, 1999). There are several examples of between-population behavioural 42 43 divergence. For instance, the effect of varying predation pressure on behaviour has been proven in common frog (Rana temporaria) tadpoles (Van Buskirk & Arioli, 2005), nine-44 spined stickleback (Pungitius pungitius) (Herczeg, Gonda, & Merilä, 2009a) or guppy 45 46 (Poecilia reticulata) (Magurran & Seghers, 1991, 1994). Such environmentally driven population divergence in behaviour has been proven to be genetically based in several species 47 (see Breden, Scott, & Michel, 1987; Brown, Burgess, & Braithwaite, 2007; Herczeg, Ab 48 49 Ghani, & Merilä, 2013; Laine et al., 2014).

50

Aggregation (gathering together) is a basic behaviour exhibited by many animals (Parrish & 51 52 Hamner, 1997; Parrish & Edelstein-Keshet, 1999), nevertheless, it does not necessarily imply any social organization (e.g. antelopes gather for water; Allaby, 1994). Aggregation can be 53 54 seen as a form of evolutionary adaptation with multiple advantages (Pitcher, 1986): it might help terrestrial species avoiding desiccation (Brockett & Hassall, 2005), it could improve 55 feeding efficiency (Heupel & Simpfendorfer, 2005) and it could provide a mechanism of 56 57 defence from predators (Broly et al., 2013). However, aggregation behaviour may also have costs, such as higher vulnerability to contact-spread parasites, increased competition for 58 resources and conspicuousness of the group towards predators (Pitcher, 1986; Romey, 1995). 59 Hence, decreased aggregation is expected in populations where the benefits of the behaviour 60 are low. For instance, the negative covariance between predation pressure and the social costs 61

of aggregation was proven experimentally (Herczeg, Gonda, & Merilä, 2009b). While many
studies were focusing on aggregation behaviour (schooling, shoaling, flocking, herding) in
vertebrates, it is also well-known in invertebrates (Hassall & Tuck, 2007; Kullmann *et al.*,
2008; Tanaka & Nishi, 2008; Broly *et al.*, 2012).

66

Sheltering (hiding under cover) is one of the most straightforward antipredator behaviours, 67 68 when it is not applied against some other environmental effects, like too harsh sunlight. Sheltering behaviour has obvious benefits, but it also incurs costs due to lost opportunities 69 connected to alternative beneficial behaviours, like mate-searching or foraging (e.g. Cooper & 70 71 Frederick, 2007; Sih, 1992). We know from previous studies that sheltering behaviour successfully reduces the risk of predation (Cooper & Frederick, 2007; Kullmann et al. 2008). 72 Further, a previous study showed that a marine isopod, *Idothea balthica*, traded shelter for 73 74 food (Vesakoski, Merilaita, & Jormalainen, 2008). The antipredatory role of aggregation and sheltering together with their predation pressure related population variation was supported in 75 76 many vertebrate taxa (e.g. fish; Magurran & Seghers, 1994; Griffiths, Armstrong, & Metcalfe, 2003; Heupel & Simpfendorfer, 2005; amphibians: Watt, Nottingham, & Young, 1997; 77 DeVito, 2003; reptiles: Gardner et al., 2016; birds: Carrascal, Alonso, & Alonso, 1990; 78 79 Forsman et al., 1998; Goodenough et al., 2017; mammals: Fryxell, 1991; Treves, Drescher, & Ingrisano, 2001). 80

81

Environmental conditions in subterranean environments differ drastically from those of the
surface. Caves and related habitats are characterized by the absence of light, food scarcity,
simplified communities and strongly buffered against daily, seasonal and yearly
environmental variation, further, as being physically separated from each other, they represent
natural replicates with no gene flow between them (Culver & Sket, 2000; Gibert &

Deharveng, 2002; Tobler et al., 2008; Culver & Pipan, 2009). Importantly, most caves are 87 88 free from both vertebrate and invertebrate predators (Culver & Pipan, 2019; White, Culver, & Pipan, 2019; but see Culver, 1975), in fact, predator avoidance was previously suggested as a 89 main factor behind cave-colonisation in several insect species (Rivera et al., 2002; Juan & 90 Emerson, 2010), as well as in Mexican tetra, Astvanax mexicanus (Romero, 1985). Further, 91 92 recent results indicate the importance of low-predation pressure behind occurrence of the 93 salamanders Eurycea lucifuga and Speleomantes strinatii in caves (Salvidio et al., 2017; Bradley & Eason, 2018; respectively). Intuitively, in populations with longer history of cave-94 adaptation, the loss or reduction of anti-predatory responses is expected. In line with this, 95 96 empirical results show that predator recognition is maintained in cave-dwelling populations of Pyrenean newt (*Calotriton asper*), a recent colonist, while such behaviour is lost in the highly 97 adapted Olm (Proteus anguinus) (Manenti et al., 2020). Despite of the consensus on the 98 99 importance of difference in predation pressure between surface and subterranean habitats, how between-population environmental variation affects behavioural traits, such as 100 101 aggregation and sheltering together is rarely tested at best. Previously, it was shown that in 102 two cave-dwelling populations of Atlantic molly, Poecilia mexicana, fish exhibit reduced shoaling behaviour (Plath & Schlupp, 2008) and the same pattern was endorsed from multiple 103 104 cave-adapted populations of A. mexicanus (Kowalko et al., 2013). However, loss of vision might be a more important factor behind such behavioural differences in these fishes than the 105 lack of predation (Kowalko et al., 2013). 106

107

Here, we studied population variation in aggregation and sheltering behaviour of common
water louse, *Asellus aquaticus*. In a common garden experiment, we tested aggregation
behaviour with and without shelter, and shelter use in general in a highly specialised cave
population (no predation, permanent darkness) and three surface populations (various

predators present, daily light fluctuations. We also applied light treatments (presence/absence) 112 113 under common garden settings. Aggregation would hardly increase foraging efficiency of a detritivorous species feeding on a wide variety of stationery food sources (e.g. living and dead 114 115 plant material, together with bacteria and fungi growing on them for surface populations, see Moore, 1975; Graça et al., 1993; Bloor, 2011; and endogenous bacterial mats in the studeied 116 117 cave, see Herczeg et al., 2020), and the potential importance of aggregating for mating was 118 ruled out by involving only males in the study. Hence, we expect that the main environmental 119 driver of the evolution of aggregation behaviour in our study system is the presence / absence of predation. The same was expected for sheltering behaviour, because sheltering in the 120 121 natural populations is unlikely to provide defence against various forms of environmental harshness, but predation. Therefore, we predicted that predation-adapted surface A. aquaticus 122 populations are more risk-averse (showing higher levels of aggregation and sheltering) than 123 124 the cave population adapted to the lack of predation. Expectations regarding light-induced plasticity are less straightforward. While vision of cave-adapted A. aquaticus is highly 125 126 reduced, they are still sensing light (see Protas, Trontelj, & Patel, 2011; Pérez-Moreno, 127 Balázs, & Bracken-Grissom, 2018; Re et al., 2018) and showing negative phototaxis (Fišer et al., 2016). Assuming that surface A. aquaticus populations are under higher predation risk 128 129 during light than at night, we predicted all populations to show higher aggregation and shelter use when tested in light, the response being the strongest in cave-adapted individuals. 130

131

132 Material and Methods

133 *Study system*

A. *aquaticus* is widespread in a wide variety of surface freshwater habitats across the Western
Palearctic (Verovnik, Sket, & Trontelj, 2005) and successfully colonized caves in Central
Europe on several independent occasions (Verovnik *et al.*, 2005; Verovnik, Prevorčnik, &

Jugovic, 2009). Similarly to other cave-dwelling species, cave adapted *A. aquaticus* exhibit
the typical troglomorphic adaptations, like reduced eyes and loss of pigmentation (PérezMoreno *et al.*, 2018; Re *et al.*, 2018).

140

Three surface populations and one cave population of A. aquaticus were used in the 141 experiments. All populations live within or in the vicinity of Budapest, Hungary. The Molnár 142 143 János Cave (47.518° N, 19.03608° E) is a water-filled cave of hydrothermal origin with a water temperature of 23-24 °C all year round. Despite the absence of physical barriers, the 144 population inhabiting the Molnár János Cave has been genetically isolated from surface 145 146 populations (including the Malom Lake population, see below) for at least 60 000 years and it shows the aforementioned troglomorphic adaptations (Pérez-Moreno et al. 2017). The only 147 available food source for this cave population of A. aquaticus is endogenous bacterial mats, 148 149 visible organic material from the surface does not enter the cave (personal observation). This latter notion is supported by results from Erőss et al. (2006) and Bodor et al. (2015), whose 150 151 results indicate no hydraulic connection between the Rózsadomb recharge area and the 152 discharge area of the Boltív Spring (connected to the Molnár János Cave - Malom lake system). The cave's outflow to the surface forms a small lake (Malom Lake) right at the cave 153 entrance (47.518° N, 19.03608° E), harbouring the first sampled surface population. Malom 154 Lake receives natural surface light, but the water temperature is similar that of the cave all 155 year round. Guppies (P. reticulata) were introduced to Malom Lake during the 20th century. 156 157 In the absence of other native fish species, guppies, forming a high-density population, are the main predators of A. aquaticus in the Malom Lake (Berczik, 1956; personal observation). The 158 remaining two surface populations, Gőtés Lake (47.59556 ° N, 19.04142° E) and Dunakeszi 159 Peat-moor (47.615613° N, 19.126392° E) are subject to natural surface light regime and 160 temperature fluctuations typical to the region. These surface populations are members of a 161

diverse community consisting competitor and predator species, *A. aquaticus* is known to be a
food source for fish and larval dragonflies in surface habitats (Harris, Karlsson Green, &
Pettersson, 2013), while the cave-dwelling population experience low biotic complexity with
a small number of competitors and absence of predators in a stable and predictable
environment.

167

168 *Collecting and housing the experimental animals*

Adult animals were collected on 30 October 2019. Samples were collected by hand sorting 169 with mash net except the subterranean animals from the Molnár János Cave where a modified 170 171 Sket-bottle was used (Chevaldonné et al., 2008) and cave diving was necessary. 2019 had an exceptionally warm autumn in the region (record 19.7 °C daily high temperature in Budapest 172 on 21 October), hence, water temperature at the surface habitats was similar to the 173 174 temperature of thermal water at Malom Lake and Molnár János Cave. After collection, animals were immediately transported to the facilities of the Biological Institute of Eötvös 175 Loránd University (Budapest, Hungary). As mentioned above, we used only males to rule out 176 the potential importance of aggregating for mating. Populations were randomly divided into 177 two subgroups and individuals were housed together in transparent plastic containers 178 179 (volume: 5 L, dimensions: $32cm \times 21cm \times 11cm$, length \times width \times height, respectively). Individuals were kept in these containers during the whole experiment (except behavioural 180 tests, see below). Water collected at the source habitats were used to fill the containers, water 181 was regularly refilled. We also provided small stones as shelters. Containers were placed in 182 custom made, light-controlled chambers (see below). Surface populations were acclimated to 183 a daily light cycle (10h light: 14h dark; controlled by a timer), while the cave-dwelling 184 population was acclimated to complete darkness and all handling processes were done under 185

red light. The temperature in the lab varied between 23-24 °C. Animals did not receive food
during the acclimation period and experiments.

188

189 Experimental setup

To video-record the animals' behaviour in different light conditions, custom made recording 190 chambers were built (100 cm \times 55 cm \times 105 cm; length \times width \times height). All chambers were 191 192 equipped with two light sources: LEDs imitating daylight (4500 K, CRI > 90) at the top and infrared LEDs (920 nm) at the bottom. This infrared wavelength is out of the visible range of 193 the animals (Dember & Richman, 2012). The chambers were closed from sides with non-194 195 transparent black plastic boards so that light did not scatter inside/ outside of the chambers. Inside each chamber, we mounted a webcam (Logitech C920 FullHD; Logitech, Lausanne, 196 Switzerland) that was modified to improve the quality of videos recorded in infrared light. 197 198 OBS Studio software (OBS Studio Contributors) was used to capture videos at 5 frames per second at HD resolution (1280 x 720). 199

200

201 We tested sheltering behaviour and aggregation in the presence of shelters on 1 November 2019. The experimental set-up consisted of circular arenas (Petri-dishes, diameter 140 mm). 202 203 The bottom of all Petri dishes was coarsened with emery paper to enable animals' normal movement (Fišer et al., 2019). These Petri dishes were housing the tested groups (see below). 204 Five shelters, made of red glass were placed into the arenas (red glass reduces light intensities 205 and filters the spectrum of light; Devigne et al., 2011). The shelters were quadratic in shape (3 206 cm wide), each of them was placed equal distances from each other inside the arenas. One 207 side of the shelter was raised slightly with a tiny piece of glass so the animals could crawl 208 under and receive thigmotactic stimuli (Fišer et al. 2019). 209

210

Animals within population were randomly divided into six groups (five individuals in each 211 212 group, sampled from the two holding tanks per population randomly). The groups were placed into small removable cylinders (diameter = 28 mm) located in the centre of the Petri-dishes. 213 214 The 24 Petri-dishes were randomly divided between two recording chambers. After ca. three minutes of acclimation, the cylinders were removed and the video-recording has started. Each 215 216 group was tested both in the presence and absence of light, treatments having different order 217 in the recording chambers. After 150 min of recording, light regimes were changed in the recording chambers and a second recording-period of 150 min started. Infrared light was on 218 for all tests, as it was needed for video recording. Aggregation in the absence of shelters were 219 220 tested on 5 November 2019. The experimental set-up was similar as in the previous experiment, except that individuals' behaviour was assessed in the absence of shelters. 221

222

223 For the video analyses, the first and the last 30 minutes of recording were used (hereafter: recording period) to see how behaviour changed along the experiment. Aggregation behaviour 224 225 in the presence of shelters was quantified by the average number of animals under the 226 occupied shelters (hereafter: 'shelter sharing') at a given observation. Sheltering behaviour was represented by the proportion of animals under shelter (hereafter 'sheltering') at a given 227 228 observation. Each behavioural variable was recorded in every third minute within these recording periods, which left us 20 observations per Petri-dish altogether (10 observations per 229 recording perios). To quantify aggregation behaviour in the absence of shelters, we measured 230 231 the distance between all pairs within the groups for every observation and then calculated the mean of these distances, thus, there was only one number per group for every observation 232 describing group cohesion (hereafter: 'distance'). For measuring the distances between 233 individuals, we used ImageJ software (Schneider, Rasband, & Eliceiri, 2012). Note that 234 individual (non-averaged) data could be analysed too, but adding the extra level of hierarchy 235

to the model would make the model problematic. However, we ran this model too and it
yielded qualitatively similar results to the analysis of averaged data, and thus we only report
the latter.

239

240 *Statistical analyses*

For analysing distance, shelter sharing and sheltering, we ran separate linear mixed models 241 242 (LMMs) by using the packages *lme4* (Bates et al., 2015) and *lmerTest* (Kuznetsova, Brockhoff, & Christensen, 2016) available in the RStudio interface 1.2.5001 (RStudio Team, 243 2020). All models were built with population, treatment (light vs. dark), recording periodt 244 245 (beginning vs. end period of the experiment) and their interactions as fixed effects. Because interpreting three-way factorial interactions is highly problematic, we included only the two-246 way interactions. Order of trial (first vs. second run within a day) was treated as a nuisance 247 248 variable and we included it in the models a single fixed effect. For the model on shelter sharing, the number of individuals hiding was also added as a fixed effect for correction. In 249 250 this model, we excluded observations where less than two individuals were hiding. Group 251 identity was also added to the models as a random effect to control for the non-independence in the data. In these models, fixed effects were tested by Wald's chi-square tests and random 252 253 effects by likelihood ratio tests. Sheltering (proportion data) was arcsine transformed (following Sokal & Rohlf, 2012) to stabilize variances. We also report the proportion of 254 explained variance by the fixed factors (marginal R^2) and by both fixed and random factors 255 (conditional R^2) available in the *MuMIn* package (Barton, 2009). 256

257

258 **Results**

The LMM on distance revealed significant population × treatment and population × recording
period interactions (Table 1). However, the only strong pattern was among populations:

individuals from the Molnár János Cave and Gőtés Lake aggregated significantly less than 261 individuals from Malom Lake and Dunakeszi Moor (Fig 1 a, b). The population × treatment 262 interaction patterns were rather weak and inconsistent, suggesting that the cave population 263 264 aggregates more in light, while the surface populations aggregate more in dark than in light. The population \times recording period interaction showed weak trends suggesting that 265 populations with higher aggregation increased, while populations with lower aggregation 266 decreased aggregation along the observation. Order of trial was also significant (Table 1), 267 268 individuals aggregated less in the second round of the experiment than in the first (data not shown). The fixed effects explained 23.4% of the total variance, while the full models 269 270 explained 49% which can be seen as sufficient explanatory power for behavioural variables. For the non-significant effects see Table 1. 271

Table 1. Result of Linear Mixed Model on aggregation behaviour in the absence of shelters in *Asellus aquaticus*. Significant effects are in bold font.

Model term	χ^2 (df)	Р
Fixed effects		
population	510.17 (4)	< 0.001
treatment	1.04 (1)	0.31
recording period	1.78 (1)	0.18
order of trial	4.22 (1)	0.04
population × treatment	18.52 (1)	< 0.001
population \times recording period	12.83 (3)	0.005
treatment \times recording period	0.43 (1)	0.51
Random effects		
Group	269.27 (1)	< 0.001







278 first vs. second recording period. Least square means \pm standard errors are shown.



 \times recording period interactions (Table 2). The population \times treatment interaction is based on

Malom Lake animals showing increased aggregation as a response to light, Gőtés Lake 282 individuals showing a weaker opposite response, while the remaining populations showing no 283 response (Fig. 2a). The population \times recording period interaction revealed that all but the 284 Dunakeszi moor individuals tended to increase aggregation along the experiment, Molnár 285 János Cave individuals showing the strongest response (Fig. 2b). Number of individuals 286 hiding had the obvious effect: when more individuals were under shelter, shelter sharing 287 increased (data not shown). The fixed effects explained 14.6% of the total variance, while the 288 289 full models explained 21.3% which can be seen as sufficient explanatory power for behavioural variables. For the non-significant effects see Table 2. 290

Table 2. Result of Linear Mixed Model on aggregation behaviour under shelters in *Asellus aquaticus*. Significant effects are in bold font.

Model term	χ^2 (df)	Р
Fixed effects		
population	57.55 (4)	< 0.001
treatment	0.78 (1)	0.38
recording period	6.03 (1)	0.01
order of trial	2.29 (1)	0.13
number of individuals hiding	6.27 (1)	0.01
population × treatment	11.79 (3)	0.008
population × recording period	13.56 (3)	0.004
treatment \times recording period	0.31 (1)	0.58
Random effects		
Group	6.24 (1)	0.01





Fig.2. Aggregation under shelters in the four tested populations of *Asellus aquaticus*. a)
aggregation in the in the presence vs. absence of light. b) aggregation in the first vs. second
recording period. Least square means ± standard errors are shown.

299

300 LMM on sheltering revealed significant population \times treatment, population \times recording period

and treatment \times recording period interactions (Table 3). The population \times treatment

302 interaction showed that all populations tended to shelter more in the light treatment than in the

dark treatment, but this reaction was particularly strong in the Malom Lake population (Fig. 303 3a). The population \times recording period interaction revealed that sheltering was similar across 304 the four populations at the beginning of the experiment, while towards the end of the 305 experiment surface populations sheltered more and the cave population sheltered less (Fig. 306 3b). The treatment \times recording period interaction indicated that the increase in sheltering 307 along the experiment was stronger in the light, than in the dark treatment (data not shown). 308 The population trends were also clear, cave-adapted individuals sheltered less than their 309 310 surface conspecifics (Fig. 3a, b). Order of trial was also significant (Table 3), individuals sheltered more in the second round of the experiment than in the first (data not shown). The 311 fixed effects explained 23.3% of the total variance, while the full models explained 34.6% 312 which can be seen as sufficient explanatory power for behavioural variables. For the non-313 significant effects see Table 3. 314

- **Table 3.** Result of Linear Mixed Model on sheltering behaviour in *Asellus aquaticus*.
- 317 Significant effects are in bold font.

Model term	χ^2 (df)	Р
Fixed effects		
population	278.31 (4)	< 0.001
treatment	81.88 (1)	< 0.001
recording period	42.44 (1)	< 0.001
order of trial	16.92 (1)	< 0.001
population × treatment	33.00 (3)	< 0.001
population × recording period	69.56 (3)	< 0.001
treatment $ imes$ recording period	9.18 (1)	< 0.001
Random effects		
Group	87.65 (1)	< 0.001



Fig.3. Sheltering in the four tested populations of *Asellus aquaticus*. a) sheltering in the
presence vs. absence of light. b) sheltering in the first vs. second recording period. Least
square means ± standard errors are shown.

324 **Discussion**

325 Aggregation and sheltering behaviours could be seen as two potential forms of evolutionary adaptation that gives animals multiple advantages, including increased defence against 326 327 predators. In the present study, we tested hypotheses about how the predator-free cave environment select for decreased aggregation and sheltering, and how cave-adaptation affects 328 329 behavioural responses to changing light conditions in the widely distributed habitat generalist 330 A. aquaticus, a small freshwater isopod that successfully colonises caves. We tested these hypotheses in a manipulative common garden experiment based on three surface and one 331 cave-adapted (showing troglomorphic phenotype and being genetically isolated from surface 332 333 populations at least 60,000 years ago; Pérez-Moreno et al., 2017) populations. The results are mixed, in some cases supporting our predictions, in others rejecting it, while some unexpected 334 patterns got also revealed. 335

336

337 Aggregation

Contrary to our initial prediction, we found no systematic differences in aggregation 338 behaviour among surface- and cave-dwelling A. aquaticus in an environment without shelters. 339 However, we found high between-population variation, two surface populations showing 340 341 considerably higher tendency for aggregation than the cave population and the third surface population (Gőtés Lake). It is possible that the Gőtés Lake population experiences lower than 342 expected predation pressure (we have no detailed fauna list with density estimates from the 343 344 studied populations), or some other environmental factor is responsible for the population variation. For instance, it is possible that in Gőtés Lake, more aggressive male A. aquaticus 345 346 are favoured than in the other surface populations, resulting in the reported patterns. Note that aggression and sociability are different personality traits (sensu Réale et al., 2007), hence, 347 they can potentially evolve independently. Aggregation behaviour might also be connected to 348

other functions than predator avoidance. For instance, aggregation could reduce the time spent 349 350 on finding mating partners and provide increased number of mating opportunities (Beauche & Richard 2013) and thus this behaviour might be affected by local population densities or 351 352 variation in operative sex ratios. According to a previous study, aggregation might be related to feeding behaviour (Heupel & Simpfendorfer 2005). However, based on our knowledge, A. 353 aquaticus is a detritivorous species feeding on various living and dead plant material in its 354 355 surface habitats, and on endogenous bacterial mats in Molnár János Cave (Herczeg et al., 2020 and references therein), thus, it is improbable that aggregation yields any foraging 356 benefits for the species. 357

358

Animals are not only aggregating in the open or during activity but they also often aggregate 359 under shelters, during inactivity (Devigne et al., 2011). In a previous study on common rough 360 361 woodlouse (Porcellio scaber Latreille, 1804), where aggregation under shelters is generally observed and individuals are showing strong thigmotaxis, researchers found the same 362 dynamics of aggregation in the presence or absence of shelters (Devigne et al. 2011; Broly et 363 al. 2012). It has also been experimentally shown that when groups of rough woodlouse in 364 bright arenas were offered two dark shelters, almost 80% of individuals in a group aggregated 365 366 under one shelter (Broly & Devigne, 2011; Broly et al. 2012). Hence, shelter use is often nonrandom, individuals can prefer shelters that are already occupied, choosing shelters with 367 larger aggregations. We expected similar patterns regarding aggregation under shelters than 368 369 without shelter, assuming that aggregation under shelters might have a relevance against predators. However, there were no clear population trends between the cave population and 370 the surface populations. 371

372

Regarding the light treatment, considering the (i) night activity of surface A. aquaticus 373 374 (Andrikovics, 1981), (ii) expected higher predation pressure during daylight for the surface populations and (iii) the negative phototaxis of the cave population, we predicted higher 375 376 aggregation in the light than in the dark treatment, the effect being the strongest in the cave population. When we tested for aggregation without shelters, the results contradicted the 377 prediction. We note that despite the significant population \times light treatment interaction, the 378 379 actual patterns seem somewhat weak. While the cave population aggregated more in light than in dark, the surface populations rather tended to aggregate more in dark than in light or 380 showed no response. Hence, there are some indirect signs of the expected negative phototaxis 381 382 in the cave population. However, the surface populations behaved in an unexpected way, and we can only speculate about the reasons. For instance, it is plausible that some social activity 383 might take place in darkness. Surface A. aquaticus is intuitively assumed to have diurnal 384 385 activity and they are indeed active during the light period. However, Andrikovics (1981) showed that the trappability of the species is three times higher during night than during 386 daylight when using passive traps, suggesting higher movement activity during night. 387 Previous experiments of ours reproduced these results under laboratory settings 388 (unpublished). Hence, the species seems to show activity all around the clock. However, even 389 390 though it is possible that some sort of increased social activity results in higher aggregations during night, but such activity among male individuals is yet unknown. Another potential 391 explanation is increased male-male aggression during the day, but again, we have no data 392 393 about it. Finally, it is possible that predation pressure is – contrary to our expectations – is higher during the night than during daylight. However, there is no data about the relative 394 395 changes in predation pressure during a day. To answer this question, future research is to be done on the species' ecology and behaviour in the wild. 396

397

In the case of aggregation under shelters, there was a weak trend for stronger aggregation in 398 the dark in Gőtés Lake, while we found a strong pattern only in Malom Lake, where 399 individuals aggregated under the shelters much stronger in the light than in the dark treatment. 400 401 During the 20th century, guppies (non-native in Hungary) were introduced to the Malom Lake, and to our knowledge, there are no native fish species present and we never observed 402 403 large insect predators (e.g. dragonfly larvae, Dytiscidae, etc.) in the lake. As a consequence, 404 guppies are under negligible predatory risk and their density in the lake is high. We repeatedly observed groups of guppies feeding on A. aquaticus. As guppies are diurnal, A. aquaticus 405 might suffer extremely higher predation pressure during the day, which explains the increased 406 407 aggregation observed in the light treatment.

408

409 *Sheltering*

410 We predicted that the cave-adapted population, evolving under the lack of predation, will use shelters less than the surface populations under presumably different, but definitely significant 411 412 predation risk. Previously, Fišer et al. (2019) showed that shelter-seeking behaviour exists in 413 A. aquaticus, but their results about differences between cave and surface populations were inconsistent, as only one cave population showed the expected decrease in shelter use. Here, 414 415 the prediction was supported, as A. aquaticus from the Molnár János Cave sheltered less than the surface populations. Obviously, we studied only one cave population, hence the results 416 cannot be generalised, but it shows that the reduced shelter use in one cave population in Fišer 417 418 et al.'s (2019) study is not an exception.

419

We also predicted, based on similar grounds that is explained in the previous subchapter, that (i) shelter use will be higher under light than under dark conditions and (ii) the difference will be more pronounced in the cave than in the surface populations. Our findings support the first prediction, but not the second. All populations sheltered more in the light treatment.
Interestingly, the reaction norms were similar in all populations but the Malom Lake, which
showed an elevated response. This pattern is congruent with our findings regarding
aggregation under shelters, where we also reported an elevated response to light in Malom
Lake. We believe that the explanation is the same: elevated shelter use and forming larger
aggregations under shelters in light are adaptations to the high predation pressure induced by
the diurnal guppies in Malom Lake.

430

431 Habituation patterns

432 All above discussed behaviours were recorded both in the beginning and at the end of the given observation period. We assumed that the beginning (being placed in the cylinder or an 433 abrupt change in light conditions) of the experiment is perceived as stressful by the focal 434 435 animals and thus they should behave differently compared to the end of the experiment after 90 minutes elapsed without disturbance. According to Blumstein (2016), 'Habituation is a 436 process that leads to decreased responsiveness to a stimulus with repeated presentation and is 437 often adaptive in that it makes it less likely that individuals will respond to harmless stimuli'. 438 In contrast, a reverse mechanism, known as sensitization, intensifies behavioural response to 439 440 constant stimulation (Bee, 2001; Stamps, Briffa, & Biro, 2012). If we treat being in a novel environment/situation as a permanent stimulus, the behavioural change during our experiment 441 can be seen as a form of habituation (e.g. Herczeg et al., 2019). Hypothesizing that cave A. 442 443 aquaticus adapted to the lack of predation will be less sensitive, we predicted that the cave population will express quicker/stronger habituation. Even though we detected a significant 444 population \times recording period interaction in all studied behaviours, the results are mixed. 445 Regarding aggregation in the absence of shelters, the significant population variation in 446 habituation revealed a weak pattern: populations with higher aggregation tended to increase, 447

while populations with lower aggregation tended to decrease aggregation along the 448 449 observation period. Whether this pattern can be seen as support for sensitisation in some and habituation in the other populations (sensu Blumstein 2016) warrants further targeted studies. 450 451 In aggregation under shelter, the Molnár János Cave population showed the strongest habituation by decreasing aggregation behaviour along the experiment. This can be seen as a 452 453 pattern supporting our prediction. However, when testing the same question in sheltering 454 behaviour, we found that the surface populations changed their behaviour more during the experiment by increasing sheltering (sensitisation), while the cave population showed only a 455 small decrease (habituation). Even though our prediction was not supported, we believe that 456 457 population variation in habituation/sensitisation governed by differences in predation pressure is an interesting idea worth pursuing in the future. 458

459

460 *Conclusions*

Taken together, we tested how does the adaptation to the predator-free, permanently dark cave 461 environment affect aggregation and sheltering behaviours and their light-induced plasticity in 462 A. aquaticus. We predicted that cave A. aquaticus will show decreased aggregation and 463 sheltering, and stronger (negative) light-induced plasticity than surface A. aquaticus. We got 464 465 mixed results. Only results about sheltering behaviour supported the predictions. However, we detected various population differences in aggregation, and light- induced plasticity in 466 general, which were unexplained by habitat type. We conclude that population divergence in 467 468 aggregation is explained by other factors than the ones differing systematically between cave vs. surface habitats. Besides further laboratory studies including more populations and also 469 470 females, understanding this system warrants extensive field surveys to reveal the relevant factors in environmental variation. More behavioural tests will be also needed for the 471 separation of the roles of sociability and aggression in the variation of aggregation behaviour. 472

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482	
483	Data availability
484	Data will be deposited in Figshare upon publication
485	
486	Conflict of interest
487	The authors declare no conflict of interest
488	
489	Author contributions
490	All authors designed the study; GeH, SS and GB collected the data; GeH and SS analysed the
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492	contribution from SS and GH; all authors reviewed the manuscript and gave final approval for
493	publication
494	
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