1	Parasite transmission in aquatic ecosystems under climate change: joint effects of
2	temperature, host behavior and elimination of parasite larvae by predators.
3	Running head: Parasite transmission and climate change
4	Gopko M. ^{1*†} , Mironova E. ^{2†} , Pasternak A ³ ., Mikheev V. ¹ and J. Taskinen ⁴
5	¹ Severtsov Institute of Ecology and Evolution RAS, Laboratory for Behaviour of Lower
6	Vertebrates
7	Moscow, Russia
8	² Severtsov Institute of Ecology and Evolution RAS, Center of Parasitology
9	Moscow, Russia
10	³ Shirshov Institute of Oceanology RAS, Plankton ecology laboratory
11	Moscow, Russia
12	⁴ Jyväskylän Yliopisto, Department of Biological and Environmental Science
13	Jyväskylä. Finland
14	* Corresponding author. E-mail: gopkomv@gmail.com, tel.: +7 (495) 954 75 53.
15	[†] Equal contribution.
16	Keywords: host-parasite interactions, disease transmission, fish behavior, predation on cercariae,
17	Diplostomum pseudospathaceum, rainbow trout, freshwater mussels, global warming
18	Abstract
19	1. A moderate raise in temperature was suggested to enhance the impact of parasites on aquatic
20	ecosystems. Under higher temperatures, poikilothermic animals (e.g. fish), increase their activity,
21	which can result in a more frequent encounter with parasites. However, temperature increase may
22	also trigger processes counteracting an increased risk of parasitic infections. For instance, removal

23 of free-living stages of parasites by filter-feeding organisms can increase with temperature and

24 potentially mitigate disease risk in ecosystems under climate change.

25 2. In our study, we aimed to find out whether an increased infection transmission under higher 26 temperatures can be, at least, partly compensated by the increased removal of parasitic larvae be 27 aquatic predators. In addition, we planned to reveal the behavioral mechanism underlying the more 28 successful transmission of the parasite at higher temperatures.

29 3.We experimentally studied how temperature, the behavior of fish host (rainbow trout) and the

30 presence of filter-feeding mussels in the environment influence transmission success of trematode

31 larvae (Diplostomum pseudospathaceum cercariae) to fish host.

32 4. We found that temperature raise increased, while presence of filter-feeding mussels in the 33 environment decreased infection intensities in fish. However, the effect of mussel's presence was 34 constant within the tested range of water temperatures (15-23°C), which suggests that it cannot 35 compensate for the observed increased transmission of parasites under temperature raise. The 36 difference in fish individual behavior (swimming activity) before the exposure to parasites was a 37 substantial factor the affecting host's vulnerability to infection. However, fish motor activity only 38 weakly correlated with temperature, therefore, it is unlikely to be responsible for the increased 39 infection success under warmer conditions. After exposure to parasites, fish strongly decreased their 40 activity. This decrease was temperature-dependent and more pronounced in bolder (more active) 41 fish, which leads to lower variability in activity of fish exposed to parasites compared with the safe 42 environment. Post-exposure activity did not influence the infection intensity.

43 5. In general, we showed that the elimination of trematode larvae by filter-feeders is unlikely to
44 deter the potential effects of global warming on host-parasite interactions in temperate freshwater
45 ecosystems.

46

2

47

48 Introduction

49 Recent studies suggest that the impact of parasites on aquatic ecosystems can be considerably 50 affected by climate change (Studer et al., 2010; Lõhmus & and Björklund, 2015; Marcogliese, 2016; 51 Cable et al., 2017). In general, though it differs from one host-parasite system to another, a 52 moderate increase in water temperature can enhance transmission of the majority of parasitic 53 species, e.g. by increasing the rate and by lengthening of the annual period of larval production, 54 affect life cycles and the global distribution of parasites (Harvell et al., 2002; Utaaker and 55 Robertson, 2015; Lõhmus & Björklund, 2015; Barber et al., 2016; Baker et al., 2018; Mouritsen et 56 al., 2018 \square . In addition, global warming causes multiple shifts in biology (growth, behavior, 57 abundance, diversity, etc.) of their hosts and predators influencing interactions of these organisms 58 with parasites (Macnab & Barber, 2011; Lõhmus & Björklund, 2015; Brunner & Eizaguirre, 59 2016). Along with the fish-host immunity suppression caused by the temperature raise (Dittmar et 60 al., 2014), changes in the behavior of receptive hosts is one of the potential mechanisms providing 61 increased parasite transmission. For instance, increased activity or ventilation rate (Pritchard et al., 62 2001; Mikheev et al., 2014; Lõhmus & Björklund, 2015) can lead to increased exposure of fish to 63 parasites.

However, temperature raise may also launch ecosystem processes, which compensate for the increased transmission success of parasites. Free-living stages of parasites comprise a substantial share of the biomass in aquatic ecosystems (Lafferty et al., 2008; Kuris et al., 2008) and many aquatic animals consume parasitic larvae, thus, significantly reducing transmission of parasites (Thieltges et al., 2008; Johnson et al., 2010; Welsh et al., 2014; Gopko et al., 2017). Feeding rates of ectothermic organisms are strongly temperature-dependent like most metabolic processes (Schmidt-Nielsen, 1997). For instance, removal of free-living stages of parasites by filter-feeders is suggested

to increase with temperature up to a threshold level determined by physiological characteristics of predators (Burge et al., 2016) \Box . However experimental data about the effect of temperature on the elimination of parasites by aquatic predators are still scarce (Goedknegt et al., 2015) \Box and do not include observations of host behavior. To our knowledge, there is only one study which reported that the presence of predators (barnacles) at higher temperatures has a stronger effect on infection transmission than at lower ones (Goedknegt et al., 2015) \Box .

77 Change in fish vulnerability to infection caused by temperature raise could be mediated by 78 fish behavior. For instance, under higher temperatures fish can increase their motor or/and 79 ventilation activity which potentially increases exposure rate, thus increasing parasite's chances to 80 penetrate host skin and gills (Mikheev et al., 2014). In addition, individual behavioral variation can also influence host vulnerability to infection. For instance, it was suggested that more risky and 81 82 exploratory individuals (i.e. individuals with higher motor activity) might be at a higher risk of infection compared with shyer ones (Hoverman & Searle, 2016; Buck et al., 2018) \Box . Though 83 84 correlation between animal behavior traits and parasitic load was suggested in many studies 85 (Hoverman & Searle, 2016; Barber et al., 2017; Cable et al., 2017)□, an influence of individual's 86 personality on vulnerability to infection has rarely been tested experimentally (see however 87 Koprivnikar et al. (2012) and Araujo et al. (2016) \Box).

A recent study showed that filter-feeding freshwater mussels *Anodonta anatina* can significantly reduce transmission of the fish trematode *Diplostomum pseudospathaceum* by eliminating its free-living stages, i.e. cercariae (Gopko et al., 2017)]. This parasite is very common in limnetic systems of temperate and boreal zones, infect a plethora of fishes, and can hamper fish farming (Valtonen & Gibson, 1997; Karvonen et al., 2006).

93 In the present study, we investigated the effect of temperature and mussels (A. anatina) on the 94 transmission of a common fish trematode (eye fluke, D. pseudospathaceum) with a focus on

95 potential interactions between these two factors and fish behavior.

96 Our main hypotheses were: (1) fish (*Onchorhyncus mykiss*) will be more vulnerable to parasitic 97 infection under higher temperature due to increased activity; (2) mussels can remove trematode 98 cercariae from the water in a wide range of temperatures and their impact on the reduction of the 99 infection in fish is temperature-dependent (i.e. they can at least partly compensate for increased 100 vulnerability to parasites in fish caused by a temperature raise).

101

102 Material and methods

103 Study objects

All experiments were conducted at the Konnevesi research station (University of Jyväskylä) in summer 2017. We used a common fish trematode *D. pseudospathaceum* as the parasite, rainbow trout *O. mykiss* as the host and freshwater mussels *A. anatina* as predators of cercariae.

107 The eye fluke *D. pseudospathaceum* has three hosts in its life-cycle: freshwater mollusks 108 (the first intermediate host), different fishes (the second intermediate host) and fish-eating birds as 109 definitive hosts (Valtonen & Gibson, 1997; Karvonen et al., 2006). In fish, this parasite localizes 110 in the eye lenses and decreases host fitness by impairing vision (Owen et al., 1993; Karvonen et al., 111 2004a) □ and manipulating host's behavior (Seppälä et al., 2004; Mikheev et al., 2010; Gopko et al., 112 2015, 2017a) \Box . Young-of-the-year rainbow trout were obtained from a commercial fish farm and 113 acclimated in the laboratory at least for two weeks before the experiments. At the fish farm, rainbow 114 trout were maintained in ground water and, therefore, were free of macroparasites. A. anatina 115 mussels were collected from Lake Jyväsjärvi and were acclimated at the lab for a week before the 116 experiments. Each mussel was observed to filter actively (siphons protruded) before the start of the 117 experiment. Infected pond snails Lymnaea stagnalis collected from Lake Konnevesi were used as a 118 source of *D. pseudospathaceum* cercariae. The shedding of cercariae by snails was checked visually

119 by incubation of snails in glasses with filtered lake water under the bright light for several hours.

120 Since in Finland (including Lake Konnevesi) *L. stagnalis* is typically infected with *D.* 121 *pseudospathaceum* rather than other related diplostomidae species (Louhi et al., 2010; Rellstab et 122 al., 2011), the cercariae were identified microscopically by their morphology.

123 Experimental design

124 The experiment was divided on 'tests' conducted seven times in a row (at different temperatures) so 125 that each test was started after the previous one ended. In each test, fish randomly chosen from the 126 stock maintained in the laboratory were placed individually in 26-28 white containers (30x40x25127 cm) filled with 12L of filtered lake water and were acclimated for an hour before exposure to 128 cercariae. Fish were randomly assigned to four treatments (6-7 replicates in each). During the 129 acclimation period, water in half of the containers was slowly warmed with aquarium heaters, while 130 in another half, similar heaters were placed, but switched off. In addition, in half of the containers in 131 each heating treatment, we placed live Anodonta anatina (one mussel per container), while in other 132 half closed empty shells of mussels. Empty shells and switched off heaters were placed in 133 containers to minimize the difference in fish behavior between the treatments (Gopko et al., 134 2017b]. Therefore, there were the following four treatments (6-7 fish in each): (1) containers with 135 heating and the presence of live mussel (H+M+), (2) containers with heating and the presence of 136 empty shell, i.e. 'mussel' control (H+M-), (3) containers with switched-off heaters and live mussels 137 (H-M+) and (4) with switched-off heaters and empty shells (H-M-).

Tests were started at the same time of the day (between 0:30 and 1:30 p.m) to exclude potential effects of the circadian rhythms. Each test lasted for two days (the first day – infection, the second – dissection). In three tests, the temperature in containers with heating was set close to 19.5°C (mean \pm SD = 19.6 \pm 1.59°C), while in four others it was around 22.5°C (22.6 \pm 1.48°C). In control containers, the temperature was about 15-17°C (16.0 \pm 0.70°C). These values are typical of the surface layer in Lake Konnevesi after wind mixing in summer (mean daily temperature range 11.9–

144 20.0, mean \pm SE = 16.1 \pm 1.23°C) (Kuha et al., 2016). Thus, the lowest water temperature in our 145 experiment reflected natural conditions in nearshore regions of this lake. In addition, these 146 temperatures are also similar to mean summer temperatures in temperate lakes (mean \pm SE = 147 16.8 \pm 0.52°C), which were calculated using data from 'laketemps' package (Sharma et al., 2015) 148 (see Supplement, Methods 1, for details). Therefore, temperatures in containers with heating reflect 149 moderate predictions of temperature increase $(1 - 5^{\circ}C)$ by the end of the 21st century (IPCC, 2014, 150 2014) being far from the most pessimistic and extreme predictions for the temperate lakes in the northern hemisphere (Sharma et al., 2007)151

The temperature was measured in each container before the first fish activity tracking (see below) and at the end of the experiments (after removing fish from containers), and did not change significantly during this period. Temperature values obtained during post-experimental measurement were used in the statistical analysis.

However, there was a substantial temperature variation both among controls in different tests (due to changes in the outside temperature) and among heated containers, because our heaters cannot be precisely calibrated. Therefore, in the statistical analysis, we treated temperature as a continuous predictor, while statistical models, where the temperature was considered as a factor are presented in a Supplement (Methods 4, Results).

In total, 180 fish were used in the statistical analysis (see, however, *Fish activity tracking* section), because 16 individuals were lost due to jumping out from containers, death for unknown reasons or obvious signs of sickness and therefore were excluded from the sample. Fish loss never exceeded 3 individuals per test and the resulting number of rainbow trout used in all treatments were similar ranging from 43 to 47 fish. Therefore, it is unlikely that an uneven fish loss in different treatments can influence the results of the statistical analysis.

167 Infection protocol and dissections

Fish were exposed to freshly produced *D. pseudospathaceum* cercariae obtained from five *L. stagnalis* snails less than 2 h before the exposure. The infection dose was 300 cercariae per fish, the
exposure time was two hours.

171 After each test, rainbow trout were caught and placed individually in 8L flow-through tanks for 24

172 hours to let parasites reach eye lenses of the fish. Then fish were killed with an overdose of MS222,

173 weighted and dissected. The number of D. pseudospathaceum metacercariae in the eye lenses of the

174 fish was counted using a dissection microscope ($32 \times$ magnification).

175 Fish activity tracking

We video recorded fish behavior at different temperatures from above the aquaria for 5 minutes before and after exposure to parasites (one hour after the addition of cercariae). A grid (10x10 cm) was drawn on the bottom of each test tank and activity was measured as a number of gridlines crossed by fish in a 5-minute interval. Records were analyzed blindly (i.e. investigator was unaware about the treatment to which an observed fish belonged). Cameras were switched on from outside to avoid the influence of the investigator on fish behavior.

182 Unfortunately, due to a technical problem, all videos from one of the tests from the mild heating 183 treatment were lost. In addition, several records were excluded from the sample, because some 184 containers were partly out of camera range. Therefore, activity video records were obtained only for 185 142 fish.

186 Statistical analysis

187 Influence of environmental conditions and fish weight on the infection intensity.

Linear mixed models were used to estimate the influence of temperature and presence/absence of alive mussel in the environment. The practical and widely used strategy to find out which variables should be included in the model is a step-down (backward) model selection, however, its too straightforward implementation (i.e. including too large a set of possibilities) can turn into a data-

192 dredging (Bolker, 2007, p. 277; Kuznetsova et al., 2017) \Box . Therefore, we first formulated a 193 biologically sensible model of interest, where all variables and the interaction purposefully tested in 194 our study were included, and then simplified the model using backward selection tool from the 195 'lmerTest' package (Kuznetsova et al., 2017) \Box .

196 The model was the following: log(infection intensity) ~ fish mass (covariate) + temperature 197 (covariate) + alive mussel presence/absence (factor) + temperature*alive mussel presence/absence + 198 experiment identity (random factor). Since we were interested in certain double interaction 199 (temperature*alive mussel presence/absence) we included only this double interaction in our model 200 of interest. The response variable (infection intensity, i.e. the number of D. pseudospathaceum 201 metacercariae in fish) was log-transformed to meet model assumptions. To verify that we did not 202 miss some important interactions, we also tested the model including all possible interactions using 203 a similar approach. The resulted models were identical (see results), which suggests that models 204 with higher order interactions are unlikely to explain the data substantially better than the model 205 obtained by the model of interest simplification.

To account for the influence of fish activity on its vulnerability to parasites, we used an abridged dataset, since recordings of rainbow trout behavior were not available for all fishes (see the explanation in the *Fish activity tracking* section) and, therefore, in this case, fish activity was included in the model of interest. In all other respects, the statistical analysis was similar to the described above. We created two separate sets of models for fish activity before and after the exposure to cercariae. P-values were calculated using Kenward – Roger's procedure for the approximation of degrees of freedom implemented in lmerTest package (Kuznetsova et al., 2017) \Box .

To present the results of the mixed-effect models graphically, partial regression plots were drawn (see the details in the Supplement, Methods 2).

215 Activity

A paired t-test was used to compare fish activity before and after the exposure to parasites. We used Fligner-Killeen test of homogeneity of variances, which is suggested to be robust against the departure from normality (Conover et al., 1981) to compare, whether variation in fish activity before the exposure to parasites was larger compared with variation after the exposure. The robust test was chosen, since the data on post-exposure fish activity violated the normality assumption (Shapiro – Wilk's test: W = 0.97, p-value = 0.005).

222 To check whether environmental conditions influence fish activity before and after addition of 223 cercariae in the containers, we started with linear mixed models where fish activity before the 224 exposure to parasites, fish activity after the exposure and differences in activities before and after 225 exposure served the response variables and experiment ID was a random factor. Presence of the 226 alive mussel in the container, temperature, fish mass and interactions between the variables were 227 components of the full model, which was then simplified using a backward selection. However, an 228 addition of the random factor did not appear to explain a substantial amount of variance in these 229 models (p > 0.3 in both cases). Therefore, the random effect was deleted from the models and we 230 proceeded with simple general linear models. The variable 'temperature' was centered by 231 subtracting the mean to make the estimates of regression coefficients more biologically sensible.

232 We also checked whether fish with a different baseline level of activity (i.e. activity before adding 233 cercriae) differed in their reaction to the presence of parasites in the environment (i.e. change in 234 activity after adding cercariae). More technically speaking, we regressed fish pre-exposure activity 235 versus the difference between pre- and post-exposure activity. However, the statistical evaluation of 236 such a relationship is usually complicated because of two methodological concerns known as 237 regression to the mean and mathematical coupling (Hayes, 1998; Tu & Gilthorpe, 2007) \Box . To 238 account for these problems, we used a method proposed by Tu et al. $(2005)\Box$. For certain formulas 239 and details see the Supplement (Methods 3), however, in brief, we calculated a correlation 240 coefficient between pre-exposure activity and difference between pre- and post-exposure activity.

Then, a correct null hypothesis was determined taking into account the correlation between pre- and post-exposure activity of fish and mathematical coupling. Finally, both observed and expected (null hypothesis) correlation coefficients were *z*-transformed and a difference between them was compared with 0 using the *z*-test.

The models, where the temperature was considered a categorical variable (three heating treatments) were also fitted (see the Supplement, Results). Their results were very similar to the ones presented in the main text of the article.

248 All statistical tests were performed using R (R Core team, 2018)□. A package 'lme4' (Bates et al.,

249 2015) was used to fit linear mixed models and get estimates of the regression coefficients.

250 'ggplot2' (Wickham, 2009)□ and 'sjPlot' (Ludecke, 2018)□ packages were utilized to visualize the

251 data.

252 **Results**

253 Infection intensity

Mean ($\pm SE$) fish weight constituted 7.77 \pm 0.15 g (total 180 ind.) and 7.48 \pm 0.15 g (142 ind. of the abridged "activity dataset"). Fish size did not differ between the treatments (with alive mussel vs control) both for full and abridged datasets (ANOVA: $F_{I, 178} = 0.13$, p = 0.72 and $F_{I, 140} = 0.12$, p =0.74 respectively). Mean \pm SE infection intensity was 46.8 \pm 2.41 in the full and 39.2 \pm 2.12 metacercariae per fish in the abridged dataset.

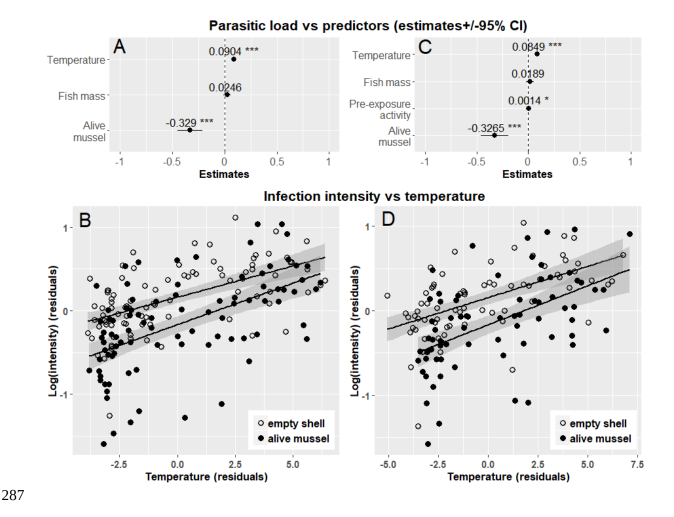
Linear mixed models comparison (i.e. procedure of backward model selection) showed that adding interaction terms did not lead to significant improvement of the model fit. Importantly, the interaction between temperature and presence of alive mussels was non-significant ($F_{1, 173.1} = 1.50$, p = 0.22, see also Table 1a), which suggests that the ability of *A. anatina* to eliminate cercariae does not change substantially under the tested temperature. Moreover, adding one of the main effects to the model (fish mass) also did not significantly increase the amount of variance explained by the

265 model (Table 1a). However, we decided to keep this predictor in the final model, since it seems 266 biologically relevant and important. When mass was excluded from the model, p-values related to 267 other predictor variables and the magnitude of estimated coefficients did not change substantially. 268 Therefore, the final model contained only the main effects and test ID (random effect) as predictors 269 (Table 1a). It showed that the effect of heating was significant and there was a 1.094-fold (exp(0.09)) 270 = 1.094) increase in parasitic load per each additional 1° C (Table 1a, Fig. 1A, C). The presence of 271 the alive mussel in the environment decreased the *D. pseudospathaceum* infection intensity in fish 272 by ~28% (Table 1a, Fig. 1A, C).

273 In the set of models, where the fish activity was included, the results were similar. Interactions were 274 also not significant and were excluded from the final model. The effect of mass was again non-275 significant, however, this predictor was left in the model for its biological relevance, as described 276 above. The effect of temperature was still highly significant (see table 1b, Fig. 1B, D). Fish activity 277 before the exposure varied substantially among fish (with range 0-278 and mean \pm SE = 117.8 \pm 4.75 278 crossed lines/5 min). The effect of fish pre-exposure activity on the infection intensity was 279 relatively week (about 1.0015 increase of infection intensity per each additional gridline crossed by 280 fish). Interestingly, when fish activity after exposure (range 1-155 and mean \pm SE = 61.4 \pm 3.21 281 lines/5 min) was added to the model instead of pre-exposure activity, it had no significant effect on 282 the infection intensity ($t_{132.2} = 0.67$, p = 0.51). The difference in fish activity before and after exposure to cercariae also was not a significant predictor of the infection intensity ($t_{132.5} = -1.77$, p =283 284 0.14).

When the temperature was added in the model as a factorial variable, the results were very similar to the presented above (see the Supplement, Fig. S1 and Table S1).

12



288 Fig. 1. Regression coefficients plots (A, C) and partial regression plots (B, D) showing the 289 influence of the temperature on the infection intensity in rainbow trout for the models fitted on the 290 full (A, B) and abridged dataset (C, D). In both cases, the presence of alive mussel in the container 291 caused a substantial decrease in the infection intensity in fish, while temperature increase led to 292 higher infection intensities. The regression lines for containers with alive mussels and control 293 containers are almost parallel, confirming the lack of interaction between the temperature and 294 presence of alive mussel in the environment. Fish, which were more active prior to the exposure, 295 were more infected compared with less active fish (A, C) (about 15% increase in the infection 296 intensity per 100 additional lines crossed by fish in five min).

 Table 1. GLMM on full and abridged datasets summary tables. Log-transformed infection

 intensity is a response variable.

		a. Full dataset models				b. Abridged dataset models					
	Fixed effects	df	t	<i>p</i> -value	Est.	SE	df	t	<i>p</i> -value	Est.	SE
	+ temperature	173.8	9.00	<0.0001	0.090	0.010	132.1	7.43	<0.0001	0.085	0.011
	+ alive mussel	173.0	-5.46	<0.0001	-0.329	0.060	132.0	-4.71	<0.0001	-0.327	0.069
	+ activity (before)						132.3	2.14	0.034	0.0014	0.0007
	+ fish mass	173.9	1.50	0.13	0.025	0.016	132.4	0.95	0.35	0.019	0.020

297

298 Activity

Paired t-test showed that before the exposure to parasites, fish were significantly more active than after the exposure ($t_{141} = 10.5$, p < 0.0001, Fig. 2A, B). Moreover, fish were more variable in activity levels before the exposure to parasites (Fligner-Killeen test: $\chi^2 = 12.43$, p = 0.0004, Fig. 2A, B). Interestingly, there was only a weak correlation between fish activity before and after exposure (Spearman's rho: $r_s = 0.19$, p = 0.03).

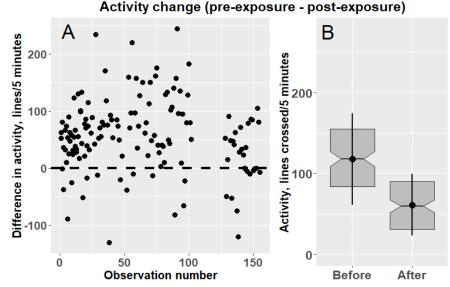


Fig. 2. Fish activity before and after the exposure to parasites. A. Difference in activity (preexposure activity minus post-exposure activity). Before the exposure to cercariae, most of the fish were more active than after the exposure (dots above the dashed zero line vs triangles below it). B. Fish activity and between individual variance in activity before the exposure was significantly higher than after it. The box on the plot represents the median with the interquartile range (IQR). The notches represent roughly 95% confidence intervals (1.58*IQR/sqrt(N)) for the medians. Dots with whiskers are mean±SD fish activity.

311 For the pre-exposure activity, we found that the model, where only the temperature was a predictor, 312 fits our data significantly better than only intercept model ($F_{1,141} = 8.49$, p = 0.004), while addition 313 of other predictors and interactions did not explain the significant additional amount of variance. 314 Fish activity increased with the temperature increase (Fig 3A) by extra four lines per each 315 additional 1°C (Estimate \pm SE = 4.27 \pm 1.47). However, when the interaction between temperature and 316 presence of alive mussel along with both main effects was included in the model its contribution 317 was marginally significant ($F_{I, 139} = 3.63$, p = 0.059, Fig. 3A). In other words, fish in both 318 treatments became more active with increasing temperature. In the presence of alive mussels, they 319 tended to increase activity even more.

320 For the post-exposure activity, the model including the presence/absence of alive mussel in the

321 container, temperature, and interactions of these effects was found the most parsimonious one.

322 There was a significant effect of temperature (Estimate \pm SE = -3.71 \pm 1.45, *t* = -2.57, *p* = 0.011) and

323 interaction between the temperature and presence of alive mussel in the model (Estimate \pm SE =

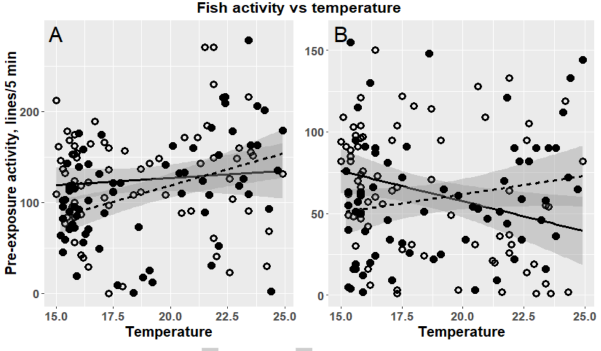
 5.95 ± 2.00 , t = 2.97, p = 0.004 Fig. 3B). It means that in the containers with alive mussels fish post-

325 exposure activity increased with temperature (regression coefficients was -3.71 + 5.95 = 2.24),

326 while in containers with empty shells fish activity even decreased with temperature raise, and the

327 slopes of the regression lines differ significantly between the treatments. Though we found a

- 328 significant influence of temperature on fish pre- and post-exposure activity in our models, the
- 329 amount of variance explained by our predictors was fairly small (6% and 7% respectively).
- 330
- 331

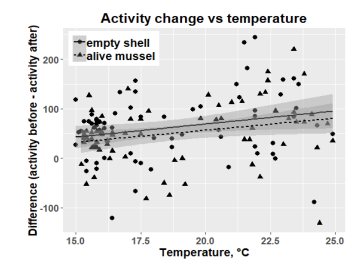


empty shell alive mussel

332 Fig. 3. Rainbow trout pre-exposure (A) and post-exposure (B) activity depending on temperature. 333 (A) Before the exposure to parasites, fish activity increased with temperature both in the presence of 334 the alive mussel and in the control (empty mussel shell) (Estimate \pm SE = 4.27 \pm 1.47 for the similar 335 slopes model). Though the slopes of regression lines for both groups did not differ significantly (p =336 0.06), the line for fish in the presence of alive mussels was steeper (dashed line). (B) After exposure, 337 the slopes of regression lines became significantly different. In the presence of alive mussels, fish 338 still increased their activity with increasing temperature, while in the control, fish decreased their 339 activity with temperature.

Temperature influenced the degree of activity change after the exposure to cercariae, i.e. preexposure activity minus post-exposure activity (Estimate \pm SE = 4.95 \pm 1.67, *t* = 2.97, p = 0.004, Fig. 4), while the addition of treatment (alive mussel/empty shell) and the interaction in the model did not increase additional amount of variance (Fig. 4). Fish changed their activity more under high temperatures compared with low temperatures.

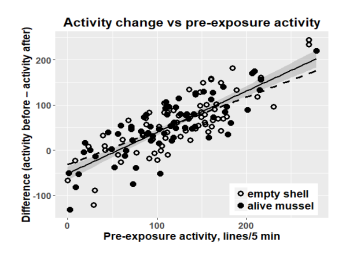
345 Fig. 4. The decrease in fish activity 346 after the exposure to parasites was 347 temperature dependent and more 348 prominent at higher than at lower 349 temperatures (extra 4.4 lines crossed 350 per each additional 1°C). However, 351 this effect was not modified by the 352 presence/absence of alive mussels 353 (regression lines are almost parallel 354 with overlapping confidence



355 intervals).

Fish more active before the exposure to parasites decreased their activity stronger compared with less active individuals (Fig. 5). The coefficient of correlation between baseline value (pre-exposure activity) and activity change was r = 0.81, which is significantly (z = 3.86, p = 0.0001) higher than null-hypothesis value (r = 0.66) calculated following Tu et al., 2005 (See Methods and Supplement).

360 Fig. 5. In both treatments fish more active 361 in the pre-exposure period decreased their 362 activity more compared with less active 363 fish (r = 0.81, z = 3.86, p = 0.0001). The 364 solid line is the empirical regression line 365 (*beta* = 0.91), while the dashed line is the 366 regression line fitted using the regression 367 coefficient (*beta* = 0.75) calculated from 368 the null-hypothetical correlation 369 coefficient value.



370 Discussion

Though temperature, predation on parasite larvae and host behavior are often reported as important regulators of parasite transmission (Lõhmus & Björklund, 2015; Barber et al., 2016, Barber et al., 2017, Burge et al., 2016, Welsh et al., 2014), their joint effect on transmission success has, to our knowledge, never been tested experimentally.

375 We found that temperature, presence of filter-feeders (freshwater mussel A. anatina) in the 376 environment and individual differences in boldness (activity in the open-field test) had a marked 377 influence on the parasite's (common eye fluke D. pseudospathaceum) infection success. Infection 378 intensity in fish increased with the temperature raise, while the presence of alive mussel led to 379 lower parasitic load in fish. Importantly, there was no interaction between these two factors, which 380 suggests that the effect of freshwater mussel on the infection transmission is constant at least in the 381 temperature range (15-23°C) tested in our study. Though the increase in the filtration rate with 382 temperature raise was demonstrated at least for several bivalve species under laboratory conditions, 383 the slope of regression curves in these studied were generally gentle (Riisgård & Seerup, 2003; 384 Kittner & Riisgård, 2005)□□. Review by Cranford et al. (2011)□□ suggested that in natural 385 conditions temperature is unlikely to be an important predictor of the feeding rate in mussels. 386 Filtration rates of mussels usually decrease under high temperatures close to the upper limit of 387 mussel's physiological tolerance (Ehrich & Harris, 2015; Burge et al., 2016)388 temperatures in our experiment were typical for natural nearshore habitats of A. anatina and did not 389 exceed comfort values for this species (Pusch et al., 2001; Falfushynska et al., 2014). Similarly to 390 our results, reduction of trematode transmission by marine bivalves (oysters) was not significantly 391 influenced by temperature, however, the hampering effect of another group of filter-feeders 392 (barnacles) increased with temperature (Goedknegt et al., 2015).

Though *D. pseudospathaceum* cercariae are known to become more infective with temperature (Lyholt & Buchmann, 1996) \Box , the mechanism of this phenomenon is unclear. One of the possible

395 explanations for it is the increase of fish motor and ventilation activity with the temperature raise 396 (Krause & Godin, 1995; Pritchard et al., 2001; Mikheev et al., 2014), which is likely to increase 397 host-parasite encounter probability (Barber et al., 2016). Our results showed that correlation 398 between fish motor activity and the temperature was surprisingly weak, however, enhanced 399 ventilation activity, which we did not measure directly, may be responsible for higher infection 400 success under increased temperatures found in our study. An alternative explanation is an immune 401 system function deterioration with temperature increase (Dittmar et al., 2014). Since our heat wave 402 was short-term, it is unlikely to have a strong influence on fish immunity, however, a performance 403 of the innate immunity providing a defense against D. pseudospathaceum infection (Scharsack & 404 Kalbe, 2014) deteriorates under warm conditions almost immediately (Dittmar et al., 2014). 405 Another possible explanation is increased activity or/and metabolism of cercariae in warmer water, 406 which can lead to a decrease in cercariae survival (Pechenik & Fried, 1995; Morley et al., 2001), 407 but at the same time can enhance parasite's infectivity (Poulin, 2006), presumably due to a short-408 term increase in parasite's host searching activity and penetration success during short time period.

409 Only fish activity before the exposure to parasites positively correlated with the parasitic load in 410 fish, while post-exposure activity did not. Fish activity before the exposure strongly varied among 411 fish and, therefore, the difference in motor activity can be a substantial factor explaining differences 412 in parasitic load among individuals. After the exposure, most of the fish decreased their activity, 413 reducing the risk of further infection (Karvonen et al., 2004b; Stumbo et al., 2012). These results 414 are dissimilar with the study on tadpoles, where no significant relationship between pre-exposure 415 activity and infection intensity was found, while activity after the exposure was negatively 416 correlated with parasitic load (Koprivnikar et al., 2012) . Previously, a decrease in fish activity 417 was reported as a possible defense against a parasitic threat (Stumbo et al., $2012)\Box$. On the other 418 hand, decrease in activity can also be a non-specific response to the presence of the alarm 419 substances released from the skin of rainbow trout (Sovová et al., 2014) damaged by penetration

420 of *D. pseudospathaceum* cercariae (Poulin et al., 2005). In general, after the exposure to parasites,

421 more active fish decreased their activity stronger than less active ones, which eventually reduced 422 variability in activity after the exposure to parasites. In non-risky conditions, bolder (more active) 423 fish may benefit from quicker food and shelter search, etc., while the main advantage of shyness is 424 lower vulnerability to new threats in the environment. Therefore, when parasite threat arises, 425 activity reduction to some optimal level may become a more beneficial strategy. This Our 426 observation is in conformity with a previous study, which showed that more bold individuals can 427 compensate risky lifestyles with a quicker and more pronounced behavioral response to the parasitic 428 threat (Klemme & Karvonen, 2016) \Box . Therefore, we suggest that the fish personality affects fish 429 vulnerability to parasites immediately after the host encounter with the parasitic threat. Later, all fish decrease their activity to a more or less uniform level. Therefore, under common environmental 430 431 threats, animal personality traits can become less expressed and behavior more uniform shrinking to 432 some optimal level. In other words, fish manifest a kind of a behavioral oddity decrease in risky 433 environments. Though oddity is a well-known factor increasing individual's susceptibility to 434 predators (Milinski, 1977, Quattrini et al., 2018, Rodgers et al., 2015), it was mainly considered 435 from the predator's point of view. However, animals' ability to self-tune their personality traits in 436 risky environments to avoid other potential threats, e.g. parasites, deserves more attention.

437 The presence of alive mussels in the environment can influence the relationship between 438 temperature and fish activity. Interestingly, in the absence of alive mussels fish post-exposure 439 activity even decreased with temperature raise, but when mussels were present in the container, the positive relationship between temperature and activity remained. It means that in a more risky 440 441 environment (without mussels, filtering cercariae) fish may try to compensate for the increased risk 442 of being infected at higher temperatures by changing their behavior in a more radical way. 443 Previously, it was shown that fish, which are less resistant against parasites can invest more in 444 developing parasites avoidance behavior compared with less vulnerable fish (Klemme & Karvonen,

445 2016).

Our research is likely to reveal only short-term ecological effects of heating within the limits of
individual plasticity of studied organisms, while global warming may cause prolonged evolutionary
processes, which also should be taken into account.

449

450 **Conclusions**

In our study, we showed that under temperature increase similar to the predicted for aquatic habitats by the end of this century, fish became more vulnerable to parasitic infection. Though filter-feeders (mussels) can effectively eliminate cercariae from the water, decreasing the parasitic load in fish, this effect remained fairly constant under a relatively broad range of temperatures. Therefore, it is unlikely that filter-feeders can compensate for the increased spread of infectious diseases with climate change as it was previously suggested (Burge et al., 2016).

457 At the first encounter with the parasitic threat, increase in vulnerability to parasites can be 458 connected with the increased activity of fish caused by the increased temperature, however, these 459 behavioral changes are unlikely to be the only factor predisposing fish to parasites under higher 460 temperatures. After more prolonged exposure to parasites, fish activity substantially decreased and 461 its influence on parasitic loads disappeared. This decrease in motor activity was temperature-462 dependent and more pronounced in bolder (more active) fish, which led to lower variability in fish 463 activity in the presence of parasites compared with the safe environment. Therefore, when working 464 together, warming and parasite threat both can influence fish behavior, altering motor activity and 465 making personality traits less expressed.

466 Acknowledgments

We want to thank the technical staff of the Konnevesi research station (University ofJyväskylä, Finland) for their assistance. The research was supported by the Academy of Finland (JT,

22

469	mobility grants 311033 and 326047); the Otto Kinne Fellowship (EM, grant 2017); the Ella and
470	Georg Ehrnrooth Foundation (MG, mobility grant 2018), the Russian Foundation for Basic
471	Research (VM, grant 17-04-00247); the Ministry of Education and Science of the Russian
472	Federation (AP, the state assignment theme 0149-2019-0008) and Russian Science Foundation grant
473	to VM and MG (19-14-00015).
474	Authors declare that they have no conflicts of interest.

475 Authors' contributions

476 All authors conceived the study. MG and EM conducted the experiments, performed the

477 statistical analysis and wrote the major part of the article. AP, VN and JT discussed the results of the

478 study, wrote minor passages of the text and revised the manuscript. JT supervised the study.

479 Data accessibility

480 All data used in the paper are stored in the figshare repository and can be accessed freely

481 (https://doi.org/10.6084/m9.figshare.8080907).

482 **References**

- 483 Araujo, A., Kirschman, L., & Warne, R. W. (2016). Behavioural phenotypes predict disease
- 484 susceptibility and infectiousness. *Biology Letters*, **12**(8), 20160480.
- 485 https://doi.org/10.1098/rsbl.2016.0480
- 486 Baker, D. M., Freeman, C. J., Wong, J. C. Y., Fogel, M. L., & Knowlton, N. (2018). Climate change
- 487 promotes parasitism in a coral symbiosis. *The ISME Journal*, **12**(3), 921–930.
- 488 https://doi.org/10.1038/s41396-018-0046-8
- 489 Barber, I., Berkhout, B. W., & Ismail, Z. (2016). Thermal Change and the Dynamics of Multi-Host
- 490 Parasite Life Cycles in Aquatic Ecosystems. *Integrative and Comparative Biology*, **56**(4), 561–
- 491 572. https://doi.org/10.1093/icb/icw025

- 492 Barber, I., Mora, A. B., Payne, E. M., Weinersmith, K. L., & Sih, A. (2017). Parasitism, personality
- 493 and cognition in fish. *Behavioural Processes*, **141**, 205–219.
- 494 https://doi.org/10.1016/j.beproc.2016.11.012
- 495 Bates, D., Mächler, M., Bolker, B., & Walker, S. (2015). Fitting Linear Mixed-Effects Models
- 496 Using lme4. Journal of Statistical Software, **67**(1), 1–48. https://doi.org/10.18637/jss.v067.i01
- 497 Bolker, B. (2007). Ecological Models and Data in R. Princeton University Press. Retrieved from
- 498 <u>https://ms.mcmaster.ca/~bolker/emdbook/book.pdf</u>
- 499 Brunner, F. S., & Eizaguirre, C. (2016). Can environmental change affect host/parasite-mediated
- 500 speciation? *Zoology*, **119**(4), 384–394. https://doi.org/10.1016/j.zool.2016.04.001
- 501 Buck, J. C., Weinstein, S. B., & Young, H. S. (2018). Evolutionary Consequences of Parasite
- 502 Avoidance. *Trends in Ecology & Evolution*, **33**(8), 619–632.
- 503 https://doi.org/10.1016/j.tree.2018.05.001
- 504 Burge, C. A., Closek, C. J., Friedman, C. S., Groner, M. L., Jenkins, C. M., Shore-Maggio, A., &
- 505 Welsh, J. E. (2016). The Use of Filter-feeders to Manage Disease in a Changing World.
- 506 Integrative and Comparative Biology, 56(4), 573–587. http://dx.doi.org/10.1093/icb/icw048
- 507 Cable, J., Barber, I., Boag, B., Ellison, A. R., Morgan, E. R., Murray, K., ... Booth, M. (2017).
- 508 Global change, parasite transmission and disease control: lessons from ecology. *Philosophical*
- 509 Transactions of the Royal Society of London. Series B, Biological Sciences, **372**(1719),
- 510 20160088. https://doi.org/10.1098/rstb.2016.0088
- 511 Conover, W. J., Johnson, M. E., & Johnson, M. M. (1981). A Comparative Study of Tests for
- 512 Homogeneity of Variances, with Applications to the Outer Continental Shelf Bidding Data.
- 513 *Technometrics*, **23**(4), 351–361. https://doi.org/10.1080/00401706.1981.10487680
- 514 Conover, W. J., Johnson, M. E., & Johnson, M. M. (1981). A Comparative Study of Tests for

515	Homogeneity of Va	ariances, with A	Applications to t	the Outer	Continental Shelf	^f Bidding Data.
010	fiomogeneity of the	and the state of t	ipplications to		Continental Shen	Diading Data

- 516 *Technometrics*, **23**(4), 351–361. https://doi.org/10.1080/00401706.1981.10487680
- 517 Dittmar, J., Janssen, H., Kuske, A., Kurtz, J., & Scharsack, J. P. (2014). Heat and immunity: an
- 518 experimental heat wave alters immune functions in three-spined sticklebacks (*Gasterosteus*
- 519 *aculeatus*). Journal of Animal Ecology, **83**(4), 744–757. https://doi.org/10.1111/1365-
- 520 2656.12175
- 521 Ehrich, M. K., & Harris, L. A. (2015). A review of existing eastern oyster filtration rate models.

522 *Ecological Modelling*, **297**, 201–212. https://doi.org/10.1016/j.ecolmodel.2014.11.023

- 523 Falfushynska, H., Gnatyshyna, L., Yurchak, I., Ivanina, A., Stoliar, O., & Sokolova, I. (2014).
- 524 Habitat pollution and thermal regime modify molecular stress responses to elevated
- 525 temperature in freshwater mussels (Anodonta anatina: Unionidae). Science of The Total

526 *Environment*, **500–501**, 339–350. https://doi.org/10.1016/j.scitotenv.2014.08.112

- 527 Fulford, R. S., Breitburg, D. L., Newell, R. I. E., & Kemp, W. M. (2007). Effects of oyster
- 528 population restoration strategies on phytoplankton biomass in Chesapeake Bay: a flexible

529 modeling approach. *Marine Ecology Progress Series*, **336**, 43–61.

- 530 Goedknegt, M. A., Welsh, J. E., Drent, J., & Thieltges, D. W. (2015). Climate change and parasite
- 531 transmission: how temperature affects parasite infectivity via predation on infective stages.
- 532 *Ecosphere*, **6**(6), 96. https://doi.org/10.1890/ES15-00016.1
- 533 Gopko, M., Mikheev, V. N., & Taskinen, J. (2015). Changes in host behaviour caused by immature
- 534 larvae of the eye fluke: evidence supporting the predation suppression hypothesis. *Behavioral*
- 535 *Ecology and Sociobiology*, **69**(10), 1723–1730. https://doi.org/10.1007/s00265-015-1984-z
- 536 Gopko, M., Mikheev, V. N., & Taskinen, J. (2017a). Deterioration of basic components of the anti-
- 537 predator behavior in fish harboring eye fluke larvae. *Behavioral Ecology and Sociobiology*,
- 538 **71**(4), 68. https://doi.org/10.1007/s00265-017-2300-x

- 539 Gopko, M., Mironova, E., Pasternak, A., Mikheev, V., & Taskinen, J. (2017b). Freshwater mussels
- 540 (Anodonta anatina) reduce transmission of a common fish trematode (eye fluke, Diplostomum
- 541 *pseudospathaceum*). *Parasitology*, **144**(14), 1971–1979.
- 542 https://doi.org/10.1017/S0031182017001421
- 543 Harvell, C. D., Mitchell, C. E., Ward, J. R., Altizer, S., Dobson, A. P., Ostfeld, R. S., & Samuel, M.
- 544 D. (2002). Climate Warming and Disease Risks for Terrestrial and Marine Biota. *Science*,
- 545 **296**(5576), 2158-2162. https://doi.org/10.1126/science.1063699
- 546 Hayes, R. J. (1998). Methods for assessing whether change depends on initial value. Statistics in
- 547 *Medicine*, **7**(9), 915–927. https://doi.org/10.1002/sim.4780070903
- 548 Hoverman, J. T., & Searle, C. L. (2016). Behavioural influences on disease risk: implications for
- 549 conservation and management. *Animal Behaviour*, **120**, 263–271.
- 550 https://doi.org/10.1016/j.anbehav.2016.05.013
- 551 IPCC, 2014: Climate Change 2014: Synthesis Report. Contribution of Working Groups I, II and III
- to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change. [Core
- 553 Writing Team, R.K. Pachauri and L.A. Meyer (eds.)]. IPCC, Geneva, Switzerland, 151 pp.
- Johnson, P. T. J., Dobson, A., Lafferty, K. D., Marcogliese, D. J., Memmott, J., Orlofske, S. A., ...
- 555 Thieltges, D. W. (2010). When parasites become prey: ecological and epidemiological
- significance of eating parasites. *Trends in Ecology & Evolution*, **25**(6), 362–371.
- 557 https://doi.org/https://doi.org/10.1016/j.tree.2010.01.005
- 558 Karvonen, A., Savolainen, M., Seppälä, O., & Valtonen, E. T. (2006a). Dynamics of Diplostomum
- *spathaceum* infection in snail hosts at a fish farm. *Parasitology Research*, **99**(4), 341–345.
- 560 https://doi.org/10.1007/s00436-006-0137-8
- 561 Karvonen, A., & Seppälä, O. (2008). Eye fluke infection and lens size reduction in fish: A
- 562 quantitative analysis. *Diseases of Aquatic Organisms*, **80**(1), 21–26.

563 https://doi.org/10.3354/dao01918

- Karvonen, A., Seppala, O., & Valtonen, E. T. (2004b). Parasite resistance and avoidance behaviour
 in preventing eye fluke infections in fish. *Parasitology*, *129*(2), 159–164. https://doi.org/DOI:
 10.1017/S0031182004005505
- 567 Kittner, C., & Riisgård, H. U. (2005). Effect of temperature on filtration rate in the mussel Mytilus
- *edulis*: no evidence for temperature compensation. *Marine Ecology Progress Series*, **305**, 147–

569 152.

- 570 Klemme, I., & Karvonen, A. (2016). Learned parasite avoidance is driven by host personality and
- 571 resistance to infection in a fish-trematode interaction. *Proceedings of the Royal Society B:*
- 572 *Biological Sciences*, **283**(1838), 20161148. https://doi.org/10.1098/rspb.2016.1148
- 573 Koprivnikar, J., Gibson, C. H., & Redfern, J. C. (2012). Infectious personalities: behavioural
- 574 syndromes and disease risk in larval amphibians. *Proceedings of the Royal Society B:*

575 *Biological Sciences*, **279**(1733), 1544–1550. https://doi.org/10.1098/rspb.2011.2156

- 576 Krause, J., & Godin, J.-G. J. (1995). Predator preferences for attacking particular prey group sizes:
- 577 consequences for predator hunting success and prey predation risk. *Animal Behaviour*, **50**(2),
- 578 465–473. https://doi.org/https://doi.org/10.1006/anbe.1995.0260
- 579 Kuha, J., Arvola, L., Hanson, P., Huotari, J., Huttula, T., Juntunen, J., ... Karjalainen, J. (2016).
- 580 Response of boreal lakes to episodic weather-induced events. *Inland Waters*, **6**(4), 523–534.
- 581 Kuris, A. M., Hechinger, R. F., Shaw, J. C., Whitney, K. L., Aguirre-Macedo, L., Boch, C. A., ...
- 582 Lafferty, K. D. (2008). Ecosystem energetic implications of parasite and free-living biomass in
- 583 three estuaries. *Nature*, **454**, 515–518. http://dx.doi.org/10.1038/nature06970
- 584 Kuznetsova, A., Brockhoff, P., & Christensen, R. (2017). ImerTest Package: Tests in Linear Mixed
- 585 Effects Models. Journal of Statistical Software, **82**(13), 1–26.

- 586 http://dx.doi.org/10.18637/jss.v082.i13
- 587 Lafferty, K. D., Allesina, S., Arim, M., Briggs, C. J., De Leo, G., Dobson, A. P., ... Thieltges, D. W.
- 588 (2008). Parasites in food webs: the ultimate missing links. *Ecology Letters*, **11**(6), 533–546.
- 589 https://doi.org/10.1111/j.1461-0248.2008.01174.x
- 590 Lõhmus, M., & Björklund, M. (2015). Climate change: what will it do to fish-parasite interactions?
- 591 *Biological Journal of the Linnean Society*, **116**(2), 397–411. https://doi.org/10.1111/bij.12584
- 592 Ludecke, D. (2018). sjPlot: Data Visualization for Statistics in Social Science.
- 593 https://doi.org/10.5281/zenodo.1308157
- 594 Lyholt, H., & Buchmann, K. (1996). Diplostomum spathaceum: effects of temperature and light on
- 595 cercarial shedding and infection of rainbow trout. *Diseases of Aquatic Organisms*, **25**(3), 169–
- 596 173.
- 597 Macnab, V., & Barber, I. (2011). Some (worms) like it hot: fish parasites grow faster in warmer
- 598 water, and alter host thermal preferences. *Global Change Biology*, **18**(5), 1540–1548.
- 599 https://doi.org/10.1111/j.1365-2486.2011.02595.x
- 600 Marcogliese, D. J. (2016). The Distribution and Abundance of Parasites in Aquatic Ecosystems in a
- 601 Changing Climate: More than Just Temperature. *Integrative and Comparative Biology*, **56**(4),
- 602 611–619. http://dx.doi.org/10.1093/icb/icw036
- 603 Mikheev, V. N., Pasternak, A. F., Taskinen, J., & Valtonen, E. T. (2010). Parasite-induced aggression
- and impaired contest ability in a fish host. *Parasites & Vectors*, **3**(1), 17.
- 605 https://doi.org/10.1186/1756-3305-3-17
- 606 Mikheev, V. N., Pasternak, A. F., Valtonen, E. T., & Taskinen, J. (2014). Increased ventilation by
- fish leads to a higher risk of parasitism. *Parasites & Vectors*, **7**(1), 281.
- 608 https://doi.org/10.1186/1756-3305-7-281

- 609 Milinski, M. (1977). Experiments on the selection by predators against spatial oddity of their prey.
- 610 *Ethology*, **43**, 311–325.
- 611 Morley, N.J., Crane, M., & Lewis, J.W. (2001). Toxicity of cadmium and zinc to Diplostomum
- 612 *spathaceum* (Trematoda: Diplostomidae) cercarial survival. *International Journal for*
- 613 *Parasitology*, **31**(11), 1211–2117.
- 614 Mouritsen, K. N., Sørensen, M. M., Poulin, R., & Fredensborg, B. L. (2018). Coastal ecosystems on
- a tipping point: Global warming and parasitism combine to alter community structure and
- 616 function. *Global Change Biology*, **24**(9), 4340–4356. https://doi.org/10.1111/gcb.14312
- 617 Owen, S. F., Barber, I., & Hart, P. J. B. (1993). Low level infection by eye fluke, Diplostomum spp.,
- 618 affects the vision of three-spined sticklebacks, *Gasterosteus aculeatus*. Journal of Fish Biology,
- 619 **42**(5), 803–806. https://doi.org/10.1111/j.1095-8649.1993.tb00387.x
- 620 Pechenik, J., & Fried, B. (1995). Effect of temperature on survival and infectivity of Echinostoma
- trivolvis cercariae: A test of the energy limitation hypothesis. *Parasitology*, **111**(3), 373-378.
- 622 doi:10.1017/S0031182000081920
- 623 Poulin, R., Marcogliese, D. J., & McLaughlin, J. D. (2005). Skin-penetrating parasites and the
- 624 release of alarm substances in juvenile rainbow trout. *Journal of Fish Biology*, **55**(1), 47–53.
- 625 https://doi.org/10.1111/j.1095-8649.1999.tb00655.x
- 626 Pritchard, V. L., Lawrence, J., Butlin, R. K., & Krause, J. (2001). Shoal choice in zebrafish, Danio
- 627 *rerio*: the influence of shoal size and activity. *Animal Behaviour*, **62**(6), 1085–1088.
- 628 https://doi.org/10.1006/anbe.2001.1858
- 629 Quattrini, F. G., Bshary, R., & Roche, D. G. (2018). Does the presence of an odd individual affect
- 630 group choice? *Behavioral Ecology*, **29**(4), 855–861. http://dx.doi.org/10.1093/beheco/ary062
- 631 Pusch, M., Siefert, J., & Walz, N. (2001). Filtration and Respiration Rates of Two Unionid Species

- and Their Impact on the Water Quality of a Lowland River. In: Bauer G., & Wächtler K. (Eds.),
- Ecology and Evolution of the Freshwater Mussels Unionoida (pp. 317–326). Heidelberg:
- 634 Springer, Berlin.
- 635 R Core team. (2018). R: A Language and Environment for Statistical Computing. Vienna, Austria.
- 636 Retrieved from https://www.r-project.org/
- 637 Rellstab, C., Louhi, K.-. R., Karvonen, A., & Jokela, J. (2011). Analysis of trematode parasite
- 638 communities in fish eye lenses by pyrosequencing of naturally pooled DNA. *Infect Genet Evol*,
- 639 **11**(6), 1276–1286. https://doi.org/10.1016/j.meegid.2011.04.018
- 640 Riisgård, H. U., & Seerup, D. F. (2003). Filtration rates in the soft clam Mya arenaria: effects of
- temperature and body size. *Sarsia*, **88**(6), 416–428.
- 642 https://doi.org/10.1080/00364820310003208
- 643 Rodgers, G. M., Downing, B., & Morrell, L. J. (2015). Prey body size mediates the predation risk
- 644 associated with being "odd." *Behavioral Ecology*, **26**(1), 242–246.
- 645 http://dx.doi.org/10.1093/beheco/aru185
- 646 Seppälä, O., Karvonen, A., & Tellervo Valtonen, E. (2004). Parasite-induced change in host
- behaviour and susceptibility to predation in an eye fluke–fish interaction. *Animal Behaviour*,
- 648 **68**(2), 257–263. https://doi.org/10.1016/j.anbehav.2003.10.021
- 649 Sharma, S., Jackson, D. A., Minns, C. K., & Shuter, B. J. (2007). Will northern fish populations be
- 650 in hot water because of climate change? *Global Change Biology*, **13**(10), 2052–2064.
- 651 <u>https://doi.org/10.1111/j.1365-2486.2007.01426.x</u>
- 652 Poulin, R. (2006). Global warming and temperature-mediated increases in cercarial emergence in
- trematode parasites. *Parasitology*, **132**(1), 143–151. https://doi.org/DOI:
- 654 10.1017/S0031182005008693

- 655 Scharsack, J., & Kalbe, M. (2014). Differences in susceptibility and immune responses of three-
- 656 spined sticklebacks (*Gasterosteus aculeatus*) from lake and river ecotypes to sequential
- 657 infections with the eye fluke *Diplostomum pseudospathaceum*. *Parasites & Vectors*, **7**(1), 109.
- 658 https://doi.org/10.1186/1756-3305-7-109
- 659 Sharma, S., Gray, D. K., Read, J. S., O'Reilly, C. M., Schneider, P., Qudrat, A., ... Woo, K. H.
- 660 (2015). A global database of lake surface temperatures collected by *in situ* and satellite
- 661 methods from 1985–2009. *Scientific Data*, **2**, 150008. https://doi.org/10.1038/sdata.2015.8
- 662 Schmidt-Nielsen, K. (1997). Animal physiology: adaptation and environment. Cambridge
- 663 University Press, Cambridge, UK.
- 664 Sovová, T., Boyle, D., Sloman, K. A., Vanegas Pérez, C., & Handy, R. D. (2014). Impaired
- behavioural response to alarm substance in rainbow trout exposed to copper nanoparticles.

666 *Aquatic Toxicology*, **152**, 195–204. https://doi.org/10.1016/j.aquatox.2014.04.003

- 667 Studer, A., Thieltges, D. W., & Poulin, R. (2010). Parasites and global warming: net effects of
- temperature on an intertidal host–parasite system. *Marine Ecology Progress Series*, **415**, 11–22.
- 669 Retrieved from https://www.int-res.com/abstracts/meps/v415/p11-22/
- 670 Stumbo, A., James, C., Goater, C., & Wisenden, B. (2012). Shoaling as an antiparasite defence in
- 671 minnows (*Pimephales promelas*) exposed to trematode cercariae. Journal of Animal Ecology,

672 **81**(6), 1319–1326. https://doi.org/10.1111/j.1365-2656.2012.02012.x

- 673 Thieltges, D. W., Jensen, K. T., & Poulin, R. (2008). The role of biotic factors in the transmission of
- 674 free-living endohelminth stages. *Parasitology*, **135**(4), 407–426.
- 675 https://doi.org/10.1017/S0031182007000248
- 676 Tu, Y.-K., Bælum, V., & Gilthorpe, M. S. (2005). The relationship between baseline value and its
- 677 change: problems in categorization and the proposal of a new method. *European Journal of*
- 678 Oral Sciences, **113**(4), 279–288. https://doi.org/10.1111/j.1600-0722.2005.00229.x

- 679 Tu, Y.-K., & Gilthorpe, M. S. (2007). Revisiting the relation between change and initial value: a
- review and evaluation. *Statistics in Medicine*, **26**(2), 443–457.
- 681 https://doi.org/10.1002/sim.2538
- 682 Utaaker, K. S., & Robertson, L. J. (2015). Climate change and foodborne transmission of parasites:
- 683 A consideration of possible interactions and impacts for selected parasites. *Food Research*
- 684 International, **68**, 16–23. https://doi.org/10.1016/j.foodres.2014.06.051
- Valtonen, E. T., & Gibson, D. I. (1997). Aspects of the biology of diplostomid metacercarial
- 686 (Digenea) populations occurring in fishes in different localities of northern Finland. Annales
- 687 *Zoologici Fennici*, **34**(1), 47–59.
- 688 Welsh, J. E., van der Meer, J., Brussaard, C. P. D., & Thieltges, D. W. (2014). Inventory of
- 689 organisms interfering with transmission of a marine trematode. *Journal of the Marine*
- 690 *Biological Association of the United Kingdom*, **94**(4), 697–702. https://doi.org/DOI:
- 691 10.1017/S0025315414000034
- 692 Wickham, H. (2009). ggplot2: Elegant Graphics for Data Analysis. New York: Springer-Verlag.
- 693 Retrieved from http://ggplot2.org