

1 **Boldness and physiological variation in round goby populations along their**  
2 **Baltic Sea invasion front**

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5 Authors: Andrea Galli<sup>a</sup> (0000-0003-0719-3135),

6 Jane W. Behrens<sup>a</sup> (0000-0002-0136-9681),

7 Manuel Gesto<sup>a,\*</sup> (0000-0002-9136-7857),

8 Nicholas P. Moran<sup>b,c</sup> (0000-0002-7331-0400)

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10 <sup>a</sup> Institute for Aquatic Resources (DTU Aqua), Technical University of Denmark, Anker  
11 Engelunds Vej 101, 2800, Kgs. Lyngby, Denmark

12 <sup>b</sup> Centre for Ocean Life - DTU Aqua, Technical University of Denmark, Kgs. Lyngby,  
13 Denmark

14 <sup>c</sup> Centre of Excellence for Biosecurity Risk Analysis, School of BioSciences, The University  
15 of Melbourne, Parkville, Australia

16

17 \*Corresponding author: Manuel Gesto (mges@aqua.dtu.dk)

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19 **Short running title:** Round goby boldness and physiological variation

20 **Abstract**

21 The round goby (*Neogobius melanostomus*) is a fish native to the Ponto-Caspian region that  
22 is highly invasive through freshwater and brackish habitats in northern Europe and North  
23 America. Individual behavioural variation appears to be an important factor in their spread,  
24 for example a round goby's personality traits can influence their dispersal tendency, which  
25 may also produce variation in the behavioural composition of populations at different points  
26 along their invasion fronts. To further analyze the drivers of behavioural variation within  
27 invasive round goby populations, we focused on two populations along the Baltic Sea  
28 invasion front with closely comparable physical and community characteristics. Specifically,  
29 this study measured personality within a novel environment and predator response context  
30 (i.e., boldness), and directly analyzed links between individuals' personality traits and their  
31 physiological characteristics and stress responses (i.e., blood cortisol and lactate, brain  
32 neurotransmitters). In contrast to previous findings, the more recently established population  
33 had similar activity levels but were less bold in response to a predator cue than the older  
34 population, which suggests that behavioural compositions within our study populations may  
35 be more driven by local environmental conditions rather than being a result of personality-  
36 biased dispersal. Furthermore, we found that both populations showed similar physiological  
37 stress responses, and there also appeared to be no detectable relationship between  
38 physiological parameters and behavioural responses to predator cues. Instead, body size and  
39 body condition were important factors influencing individual behavioural responses. Overall,  
40 our results reinforce the importance of boldness traits as a form of phenotypic variation in  
41 round goby populations in the Baltic Sea. We also highlight the importance of these traits for  
42 future studies specifically testing for effects of invasion processes on phenotypic variation in  
43 the species. Nonetheless, our results also highlight that the physiological mechanisms  
44 underpinning behavioural variation in these populations remain unclear.

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48 **Keywords:** invasive, personality, risk-taking behaviour, predation, novel environment, stress

## 49 **1. Introduction**

50 Biological invasions are multi-phase processes that can have serious impacts on invaded  
51 ecosystems, particularly in marine and estuarine environments. These ecosystems are  
52 particularly exposed to invasion due to human-driven introduction pathways such as shipping  
53 (Kotta et al., 2016), recreational boating, aquaculture and even aquarium trade (Williams &  
54 Grosholz, 2008). The round goby (*Neogobius melanostomus*, Pallas, 1814) is a highly  
55 successful invasive species in these coastal habitats. Most likely introduced from ballast  
56 water from the Ponto-Caspian region, the first appearance of round gobies within the Baltic  
57 Sea region was in the Gulf of Gdansk in Poland in early 1990 (Skóra & Stolarski, 1993). By  
58 the end of the decade, it became the dominant fish in shallow waters of the western part of  
59 the gulf (Sapota & Skóra, 2005). Since then, this predatory fish has spread west along the  
60 coasts of Poland, Germany, and now into Denmark. In 2009 the species was first observed in  
61 the inner Danish waters south of Zealand in Guldborgsund (Fig. 1), and since then it has  
62 spread on average 30 km yr<sup>-1</sup> along the coastline of Zealand and the islands in  
63 Smålandsfarvandet (Azour et al., 2015). In the Baltic Sea, round gobies appear to eat the eggs  
64 of native species, including flounder (*Platichthys flesus*), and native gobies (Karlson et al.  
65 2007), and can have major impacts on the abundances and composition of local benthic  
66 communities via their feeding behaviour (Kipp et al., 2012; Pennuto et al., 2018; van Deurs et  
67 al., 2021).

68

69 Personality is often defined as behavioural variation that shows consistent variation among-  
70 individuals over time (or context), measured as the relative proportion or component of among-  
71 individual variance estimated from repeated behavioural measurements (Dall et al., 2004;  
72 Sánchez-Tójar et al., 2022). Personality differences in boldness, aggressiveness, activity, and  
73 sociability have been linked to dispersal (Cote et al., 2010; Hirsch et al., 2017; Myles-Gonzalez  
74 et al., 2015; Rehage & Sih, 2004), and modelling has also suggested that greater diversity of  
75 behavioural traits within a population may greatly accelerate invasion rates (Elliott and Cornell  
76 2012). Personality may also influence dispersal and invasion spread at multiple stages,  
77 including decisions to stay or depart, when and where to settle, and post-dispersal success  
78 (Chapple et al., 2012; Weis & Sol, 2016). In a study about western bluebirds (*Sialia Mexicana*),  
79 Duckworth (2008) demonstrated that more aggressive individuals lead dispersion at the  
80 invasive front, and after establishment they are substituted by less aggressive individuals  
81 because of their poor parental care. These differences in personality types are spread along the  
82 invasion succession and can produce populations at different stages of invasions with

83 contrasting behavioural compositions, including round goby populations (Myles-Gonzalez et  
84 al., 2015; Thorlacius, Hellström, Finn, et al., 2015). This is likely to influence how the invasive  
85 populations at different stages of an invasion impact the communities in recipient ecosystems,  
86 as personality can be linked to habitat use and foraging behaviour in individuals (Moran, Wong,  
87 et al., 2017; Patrick & Weimerskirch, 2014). Nonetheless, there is still limited data on  
88 behavioural variation within invasive populations along invasion fronts, and what underlying  
89 mechanisms can produce these behavioural differences.

90  
91 Several underlying mechanisms or proximate causes might lead to personality variation,  
92 which may be linked to genetic/epigenetic variation, phenotypic plasticity in response to  
93 individual-level differences in environmental/state variables, or the interaction between the  
94 two (Dewitt & Scheiner, 2004; Pigliucci, 2005; Wolf & Weissing, 2010). Differences  
95 between individuals' intrinsic states can lead to differences in behaviour. For example,  
96 (Behrens et al., 2020) found that body size was associated with boldness in round gobies  
97 where bolder fish tended to be smaller, potentially due to the metabolic cost of their  
98 behaviour that results in a slower growth rate. Body condition may similarly influence  
99 behaviour (Moran et al., 2021), where lower body condition is often associated with higher  
100 risk-taking. Also, differences in hormone levels (Niemelä & Dingemans, 2018) and certain  
101 neurotransmitters (Ferris & Delville, 1994) can produce differences in behaviour.

102  
103 Stress triggers a neuroendocrine response in vertebrates that results in the production of  
104 corticosteroids and catecholamines (Wendelaar Bonga, 1997). Cortisol is the main  
105 corticosteroid in teleost fish and is widely used as a stress marker. Among other functions, it  
106 increases energy availability through gluconeogenesis, complementing the action of  
107 catecholamines mobilizing glucose from glycogen stores. This availability of energy  
108 facilitates any necessary physical responses to the stressor. Another parameter used to  
109 measure stress response in vertebrate is lactate. Lactate is a product of anaerobic metabolism,  
110 usually triggered when oxygen supply to tissues is in shortfall, for example during strenuous  
111 exercise/muscular activity. High swimming activity occurs often as part of the behavioural  
112 response to stress (Wells & Pankhurst, 1999), leading to an increase of lactic acid production  
113 and, thus, to a higher concentration of lactate in blood. Brain monoaminergic  
114 neurotransmitters such as dopamine and serotonin are believed to have a prominent role in  
115 the regulation/organization of the stress response of vertebrates, and seem to be at least  
116 partially responsible for the differences in stress response associated to personalities

117 (Crawford et al., 2010; Winberg et al., 2016). Serotonergic activity, in particular, appears  
118 consistently elevated in certain areas of the vertebrate brain during acute stress (Emerson et  
119 al., 2000; Gesto et al., 2013) and can be also affected by persistent or repeated stressors  
120 (Winberg & Thörnqvist, 2016). Importantly, both stress-response phenotypes and brain  
121 monoamines are linked to individual behavior and personality (Soares et al., 2018; Winberg  
122 & Sneddon, 2022).

123

124 The overall goal of this study is to measure differences in boldness/risk-taking behaviour and  
125 physiological stress responses of round gobies in two populations across their invasion path  
126 (Grønsund and Søvang, first recorded in 2011 and 2016, respectively) in the southeast Baltic  
127 Sea (Fig. 1). By comparing these populations, we specifically tested the following research  
128 questions:

- 129 1. *Do round gobies within these two populations show among-individual differences in*  
130 *boldness/risk-taking behaviour in a novel environment and/or predator response*  
131 *context (i.e., personality)?* We predicted that both populations would show individual  
132 differences in boldness traits, as consistent among-individual variation has commonly  
133 been found in round gobies (Myles-Gonzalez et al., 2015; Thorlacius, Hellström,  
134 Finn, et al., 2015; Thorlacius & Brodin, 2018; Behrens et al., 2020).
- 135 2. *Is there variation in boldness/risk-taking behaviour associated with (a) the*  
136 *population, and/or (b) the physical state (i.e., body size, body condition) of round*  
137 *gobies?* Some studies have shown that the populations nearer the invasive front tend  
138 to be bolder in novel environments (Myles-Gonzalez et al., 2015; Thorlacius,  
139 Hellström, Finn, et al., 2015) consistent with personality-biased dispersal favoring  
140 bolder individuals, therefore we expected to find bolder fish in the newer population  
141 (i.e. Søvang). We also expect smaller fish and fish in lower body condition to show  
142 higher levels of boldness (following Behrens et al., 2020; Moran et al., 2021).
- 143 3. *Are there (a) population differences in the physiological responses to acute stress,*  
144 *and (b) are physiological and behavioural responses linked?* The hormonal and  
145 neurochemical state of individuals may be important sources of variation underlying  
146 behavioural variation among individuals (Sih et al., 2015; Wolf & Weissing, 2010).  
147 Therefore we expect increased boldness/risk-taking to be negatively correlated with  
148 blood cortisol and lactate responses in individuals from both populations, and  
149 dopaminergic and serotonergic activity in the brain (Gesto et al., 2013). If we find

150 higher boldness/risk-taking in Søvang (as above), we also expect to find lower  
151 physiological stress responses in Søvang.

152

## 153 **2. Methods**

### 154 *2.1. Fish and holding conditions*

155 Fish were collected from two sites, Grønsund (54.90371°N, 12.10367°E, established 2011)  
156 and Søvang (55.57146°N, 12.62579°E, established 2016) (Fig. 1). Both of which are shallow  
157 brackish coastal areas, with predominantly sandy substrate with scattered boulders, where  
158 round gobies are typically found in high abundances particularly during spring and summer  
159 periods (Azour et al., 2015). Collection occurred in May 2021 as part of a broader sampling  
160 effort to collect round gobies at numerous sites along their east-coast Danish invasion front.  
161 Fish from these two sites chosen for this experiment, as they both had similarly high  
162 abundances at sampling and both showed very low mortality during laboratory acclimation,  
163 limiting survivorship bias effects on experimental fish.

164

165 Fish were caught using a combination of passive sampling methods set overnight (i.e. double  
166 funnel fyke nets, baited box and cylinder traps) to reduce the potential for personality-biased  
167 sampling (Biro & Dingemanse, 2009; Michelangeli et al., 2016). Physical conditions  
168 measured within sites during sampling were similar (Søvang 16/5/21: 8.75ppt, 14.1 °C;  
169 Grønsund 18/5/21: 8.27ppt, 13.2 °C). Fish were transported to DTU Aqua, Lyngby,  
170 Denmark, and acclimated to laboratory conditions for 48 hours before being  
171 individually tagged with passive integrated transponder (PIT) tags (12 × 2 mm, 0.1 g, Oregon  
172 RFID Inc.), using previously published methods that have minimal effects of fish health and  
173 condition (Jørgensen et al., 2017).

174

175 Before the start of this experiment, fish were held in two large mixed-sex and mixed-  
176 population round holding tanks (3000 L) enriched with artificial eel grass and connected with  
177 a recirculating water system. Fifteen days before the behavioural experiment, experimental  
178 fish (n = 36 per population) were randomly selected and allocated to three round tanks (800  
179 L) in groups of 24 per tank. Only males were selected for this experiment as both populations  
180 were heavily male-biased (Søvang, 90.2% male; Grønsund, 86.9% male at sampling), and  
181 there were too few females to include sex as a factor in analysis. The lack of females in goby  
182 samples is commonly found in invasion front populations, which may be results of male-  
183 biased spread and dispersal producing male-biased populations, and/or because the males are

184 often found to be more active, facilitating their capture with passive traps (Corkum et al.,  
185 2004; Marentette et al., 2011).

186

187 The salinity in the holding tanks was  $10.5 \pm 0.5$  ppt and the temperature was constant at  $10.5$   
188  $\pm 0.1^\circ\text{C}$  with a daily cycle of 12 hours of light and graduated increases/decreases in light  
189 intensity at dawn and dusk to simulate a natural light regime. During the whole project fish  
190 were fed every two days with dry pelleted feed (3mm Ivory Ex composite pellets, Aller  
191 Aqua, Denmark). Experimental fish were fed on the day before behavioural trials. Before the  
192 physiological sampling feeding was stopped for 48 hours before the 16 hours of isolation, to  
193 help maintain water quality in their individual holding tanks before blood and brain sampling.

194

## 195 2.2. *Experiment A: Behavioural repeatability and responses to stress*

196 After acclimating to their holding tanks over four weeks, all fish from one holding tank fish  
197 were assayed per day over 3 days, followed by repeat trials the following week. Individuals  
198 were selected using random sweeps through the holding tanks, and fish were not identified  
199 until after trials, so the order was blinded and assumed to be random within tank groups/trial  
200 days. The experiment started at 9:00am and every 30 minutes two fish were moved from the  
201 holding tank to individual black tanks (24 x 34 x 15.5 cm) covered with brown cardboard and  
202 aired with air stones for 2 h to standardize pre-trial handling and stress levels. After isolation,  
203 the fish were moved to the behavioural arenas using the individual tanks still covered and  
204 opened only once they are immersed in the arena water.

205

206 The behavioural arenas consisted of two opaque PET-plastic white tanks, A and B (internal  
207 dimensions, 32.25 x 49.25 cm), surrounded and separated by white polystyrene sheets to  
208 insulate the arenas from both external sound and visual inputs (Fig. 2). The experiment was  
209 performed under constant laboratory fluorescent lighting (approximately 45-50 lux within  
210 arenas at the water surface), with temperature and salinity identical to the holding tanks. One  
211 camera (a modified Logitech BRIO 4K Ultra HD webcam, Logitech, Switzerland) connected  
212 to a laptop (Logitech Capture, version 2.06.12) recorded the tanks from 1 m above. After a  
213 brief 5 min acclimatization, behaviour was recorded for a further 5 minutes to provide  
214 baseline behavioural measures (*pre-strike* period). A simulated bird strike was then  
215 performed using a bolt (5.5 cm, 23 g) suspended 80 cm above the water in the centre each  
216 tank by fishing lines (method adapted from Behrens et al., 2020). Bolts were released to hit  
217 the water surface and penetrate 5 cm into the water column, then were immediately retracted

218 to their original positions. Behaviour was recorded for 5 minutes after the strike before the  
219 trial was terminated (*post-strike* period). After every run the arenas were rinsed with  
220 deionized water and the water was entirely replaced using freshly filtered water from the  
221 laboratory's recirculating system, to prevent contamination from the previous fish by  
222 conspecific chemical cues that could alter the stress level during the experiment (Barcellos et  
223 al., 2011).

224

225 Videos were analyzed by using the software Toxtrac (Rodriguez et al. 2018, version 2.96).  
226 Each video/arena was individually calibrated to account for any minor variation in the  
227 positions of tanks and the camera in each trial. Behavioural variables, activity, centre area use  
228 and time spent frozen (see Table 1), were collected in both the *pre-strike* and *post-strike*  
229 periods to represent boldness/risk behaviours associated with fish movement in a novel  
230 environment and predator response context respectively (Moran et al., 2021). Direct  
231 responses to the predator strike cue were quantified as the latency to freeze following the  
232 strike, and latency to resume movement after freezing (as per (Behrens et al., 2020), both of  
233 which were quantified within ToxTrac (see details in Table 1).

234

### 235 2.3. *Experiment B: Physiological responses to stress*

236 Physiological stress response was measured in fish a minimum of 10 days after completing  
237 behavioural trials. Groups of 12 fish were moved each day from holding tanks to individual  
238 covered black tanks (24 x 34 x 15.5cm) for 16 hours to allow cortisol concentration in the  
239 blood decrease to baseline levels. The tanks were well aerated with an air stone and covered  
240 by green tarpaulin over 80% of the tank to reduce external sounds/visual inputs and provide a  
241 low stress individual holding environment for each fish.

242

243 Fish were randomly allocated to three treatment groups: 1) *sham control*, exposed to 10 mL  
244 of unscented water (n = 12 per population); 2) *cue*, exposed to 2,5 mL of water with chemical  
245 conspecific alarm cue mixed with additional 7,5 mL of unscented water (n = 18 per  
246 population); and. 3) *baseline control*, sampled after 16 hours without the addition of any  
247 stressor (n = 3 per population). The chemical alarm cues were produced to mimic chemicals  
248 released by an injured conspecific (e.g., due to a predator attack) using a method based on  
249 Smith (1989). Specifically, one non-experimental fish from laboratory stock was euthanized  
250 with an overdose of tricaine methanesulphonate ("MS-222", Acros Organics) and placed in a  
251 clan petri dish: 25 cuts were done on each flank by using a scalpel, then the fish was rinsed



252 with 15 mL of marine water. 2,5 mL of this solution were mixed with 7,5 mL of marine  
253 water, to be added to their individual holding tanks. A small hole in the tarpaulin allowed for  
254 the administration of *cue* treatments, while avoiding visible contact between the operator and  
255 the fish.

256

257 The *baseline control* group were sampled first, while the *cue* and *sham control* groups were  
258 sampled between 30 and 45 minutes (short exposure) and 75-90 minutes (long exposure) after  
259 the adding the *cue* mixtures to also allow for the analysis of exposure time on physiological  
260 responses. Each fish was then exposed to an overdose of tricaine methanesulfonate (MS-222  
261 500 mg/L), and blood samples were collected immediately using a heparinized syringe (heparin  
262 lithium salt, Sigma-Aldrich, Denmark; diluted 1 mg/mL in 0.9 % NaCl) (Marentette et al. 2013)  
263 from the caudal vein and stored on ice. Blood samples of 0.3 mL were stored in 1.5 mL  
264 Eppendorf tubes containing 15 $\mu$ L of the heparin solution. The fish was then measured for  
265 weight and total and standard length ('SL') before the head was removed with a sharp knife  
266 and placed immediately in aluminum foil within plastic sample bags on dry ice. Blood samples  
267 were then centrifuged (accuSpin<sup>TM</sup> Micro 17R, Fisherbrand<sup>TM</sup> Microcentrifuges, UK) at 2500  
268 x g for 2 minutes to obtain plasma, and both plasma and whole head samples were stored  
269 at -80°C with the heads until the analysis.

270

271 Plasma cortisol was quantified by means of a commercial ELISA kit (ref. 402710, Neogen  
272 Europe, Ayrshire, UK). Plasma lactate was analyzed with a colorimetric kit from Sigma-  
273 Aldrich (ref. MAK064, Merck, Darmstadt, Germany). Frozen fish heads were left at room  
274 temperature for 5 min and the telencephala were dissected out from fish brains while still  
275 semi-frozen, after removing the roof of the skull with a scalpel. This brain region was  
276 selected for analysis for having a key involvement in the regulation of emotional reactivity,  
277 including stress responses in fish (Vindas et al., 2018). Telencephala were weighed and then  
278 homogenized in 400  $\mu$ L of a 0.4 M perchloric acid, 0.1 mM EDTA solution, using  
279 ultrasounds (Sonopuls ultrasonic homogenizer, Bandelin, Germany). Homogenates were  
280 centrifuged (14000 x g, 4°C, 10 min), and the supernatants were immediately analyzed by  
281 means of HPLC with electrochemical detection, as described elsewhere (Alfonso et al.,  
282 2019). The levels of serotonin, dopamine, and their respective main oxidative metabolites, 5-  
283 hydroxyindoleacetic acid (5HIAA) and 3,4-dihydroxyphenylacetic acid (DOPAC) were  
284 quantified (for further details of the physiological variables used in the study, see Table 2).

285

286 2.4. *Statistical analysis*

287 All the analysis was performed in R (v4.1.2, R Core Team, 2013). Before behavioural  
288 analysis, variables were transformed where necessary (see Table 1). Intraclass correlation  
289 coefficients were used to calculate the repeatability of behavioural variables (package “rptR”,  
290 v0.9.21, Stoffel et al., 2017), which represents the proportion of among-individual variance  
291 relative to total phenotypic variance for each variable (Nakagawa & Schielzeth, 2010). To  
292 assess repeatability across all experimental fish, we calculated raw repeatabilities and  
293 population-adjusted repeatabilities. Adjusted repeatability includes population as a random  
294 effect to remove influence of population-level differences on the among-individual variance  
295 component. Closely related populations may show different levels of behavioural  
296 consistency, which may be related to local environmental conditions or dispersal processes  
297 (as in Moran et al., 2017; Thorlacius, Hellström, & Brodin, 2015), therefore, repeatability  
298 was also estimated for each behavioural variable within each population. Data for four  
299 replicate trials were lost due to technical problems with video recording and/or tracking (trial  
300 1, IDs: #447, #9772, #9888, #9955). Data for three fish were also removed from all  
301 behavioural analysis due to injuries (IDs: #485, #9765, #9924) and cataracts (#9765) found  
302 during physiological sampling, as injuries incurred during the experiment may affect their  
303 behavioural responses and cataracts are known to influence predator responses in this species  
304 (Flink et al. 2017).

305

306 Effects of population and condition on behavioural responses to the predator cue were tested  
307 using linear and generalized mixed-effect models (package “lme4”, v1.1-27.1, Bates et al.  
308 2015). Behavioral variables included here were those that showed significant non-zero  
309 repeatability (i.e., activity, centre area use and freeze time) so represent measures of among-  
310 individual behavioural (personality) variation in these populations (see Table 3). To test for  
311 changes in behaviour in response to the predator cue, behaviour from the *pre-* and *post-strike*  
312 ‘period’ were analyzed together, where the interaction between fixed effects and period  
313 represents its effect on the behavioural response of fish to the predator cue. Fixed effects, fish  
314 length (SL), body condition, population, and replicate number (and their interaction with  
315 period) were included. There appeared to be a correlation between SL and Fulton’s condition  
316 factor ( $r_{\text{pearson}} = -0.25$ ,  $p = 0.036$ ) (Ricker, 1975), therefore a group-specific condition factor  
317 was calculated using the slope of a  $\log_e(\text{SL}) - \log_e(\text{weight})$  regression line (based on Green,  
318 2001). To confirm that weight-length relationship is similar across sites, we tested for a  
319  $\text{site} * \log_e(\text{weight})$  interaction. This interaction was not significant ( $p\text{-value} = 0.519$ ), so a

320 common slope was used. Condition factor (' $K$ ') was calculated as:  $K = W / SL^n * 100$ , where  
321  $W$  is the weight (g),  $SL$  is the total length (cm) and  $n$  is the slope of the joint regression line  
322 (i.e. 2.7468). Continuous fixed effect factors (i.e.  $SL$ ,  $K$ , and replicate number) were then Z-  
323 scaled for analysis to improve the interpretability of regression coefficients within models  
324 (Schielzeth, 2010). In addition to fish ID, random effects initially included trial arena,  
325 round/order, and holding tank as potential grouping factors/sources of non-independence  
326 within the trials, although arena and holding tank were removed from all final models as they  
327 explained very little or no variance. Conditional and marginal  $R^2$  values were also calculated  
328 to estimate to total proportion of variance explained by fixed and random effects in the  
329 models, and the proportion of variance explained by fixed effects only, respectively (package  
330 'performance', v0.7.0, Lüdtke et al., 2021).

331

332 Linear mixed-effect models were used to test for effects on physiological response variables  
333 (Table 5). Response variables were log transformed to approximate normality and the  
334 continuous fixed effect factors were also scaled as above. Fixed effects included body size  
335 ( $SL$ ), condition factor, exposure time (i.e., as short versus long cue exposure), population and  
336 treatment (*sham control* v *cue*). A population\*treatment interaction was also included to  
337 specifically test if populations responded differently to the alarm cue. Additional interactions  
338 (as in behavioural models above) were not included here to limit overparameterization of the  
339 models, particularly given the smaller number of replicates per treatment and additional  
340 factors being analyzed. The *baseline control* group was used to provide a qualitative  
341 comparison with other treatment groups only so was not analyzed in these models. Random  
342 effects initially included were holding tank (i.e. tank A, B, C) and sampling day (i.e. A1, A2,  
343 B1, B2, C1, C2), to account for these potential sources on non-independence, but were  
344 removed if they resolved little or no variance. As above, data from injured fish were removed  
345 from analysis, and samples for two additional fish could not be included due to technical  
346 issues with sample collection and processing (#9928, #9822).

347

348 For fish that were included in both behavioural and physiological analysis, we tested for  
349 correlations between physiological responses (in *cue* treatments group only) and behavioural  
350 variables (*pre-* and *post-strike* separately). Ideally, repeated measures of physiological and  
351 behavioural variables would allow us to analyse co-variance across both within and among-  
352 individual levels (Dingemanse and Dochtermann 2012), although this is not possible using

353 these physiological sampling methods. Therefore, we have calculated correlations between  
354 physiological variables at the phenotypic level, and among-individual behavioural variation  
355 (i.e. by using an individual's average behavioural score across trials). Populations were  
356 analyzed separately. Spearman's rank (non-parametric) correlations were used with  
357 untransformed response variables. We present correlations without adjustments for multiple  
358 comparisons, so the significance of any one single significant correlation should be  
359 interpreted cautiously, although this was not an issue with this dataset.

360

### 361 2.5. *Animal ethics statement*

362 Ethical permit 2017-15-0201-01282 from the Danish Animal Ethics Committee  
363 (Dyreforsøgstilsynet) and its extensions covered all experiments reported here.

364

## 365 3. Results

366 Activity, centre use and freezing time variables showed significantly non-zero repeatability  
367 associated with fish ID between behavioural trials (Table 3), and estimates were largely  
368 similar in the *pre-strike* and *post-strike* periods. This suggests behavioural variation among  
369 individuals is maintained under acute predation pressure. Behaviour was also highly  
370 correlated between *pre-* and *post-strike* periods for each of these variables ( $\text{activity}_{\text{pre-post}}$ ,  
371  $r_{\text{spearman}} = 0.72$ ,  $p < 0.001$ ,  $\text{centre use}_{\text{pre-post}}$ ,  $r_{\text{spearman}} = 0.56$ ,  $p < 0.001$ ,  $\text{freeze time}_{\text{pre-post}}$ ,  
372  $r_{\text{spearman}} = 0.73$ ,  $p < 0.001$ ). In contrast, both latency to freeze and latency to recover did not  
373 show significant repeatability associated with fish ID. Within populations, activity and  
374 freezing time showed significant repeatability in the older population (Grønsund, est. 2011)  
375 but not the newer population (Søvang, est. 2016) in both periods, whereas for centre use  
376 Søvang fish showed strong repeatability relative to Grønsund fish (see Table 3).

377

378 Populations showed different activity-level responses to the predator strike (i.e.,  
379  $\text{period} \times \text{population}$  interaction effect), where Søvang fish reduced their activity more in the  
380 *post-strike* period relative to the Grønsund fish (Table 4, Fig. 3a). There was no significant  
381 population effect on activity, only a trend showing that Søvang fish were slightly less active  
382 (see Fig 3a). Instead, body size (SL) was a strong predictor of activity level, where smaller  
383 fish appeared more active overall, but showed greater reductions in their activity following  
384 the predator strike relative to larger fish (i.e.,  $\text{length}$  and  $\text{period} \times \text{length}$  effects, Table 4, Fig.  
385 3b). Therefore, the lack of population difference despite the apparent trend of lower activity  
386 in Søvang fish may have been a result of their slightly larger average body size than

387 Grønsund fish [ $SL_{Grønsund} = 13.10$  (s.d. 1.54);  $SL_{Søvang} = 14.61$  cm (s.d. 1.61); for further  
388 details see supplementary materials S1, Fig. S1].

389

390 Fish overall reduced their use of centre areas following the predator strike (i.e., period effect,  
391 Table 4), and both condition and length influenced individual responses, with larger fish and  
392 fish in better body condition being less responsive to the predator strike (i.e., period\*length,  
393 period\*condition effects, Table 4). Population had no effects on centre use. Body size (SL)  
394 also showed effects on freezing time, where smaller fish spent less time frozen overall, but  
395 increased their time spent frozen following the predator strike more relative to larger fish  
396 (i.e., length and period\*length effects respectively, Table 4). There were no effects of  
397 condition or population on freezing time. Note, the three repeatable variables were all  
398 strongly correlated to each other despite showing differing effects from length, condition, and  
399 population (activity-centre use,  $r_{\text{spearman}} = 0.57$ ,  $p < 0.001$ ; activity-freezing time,  $r_{\text{spearman}} = -$   
400  $0.95$ ,  $p < 0.001$ ; centre use-freezing time,  $r_{\text{spearman}} = -0.56$ ,  $p < 0.001$ ), such that more active  
401 individuals also spent less time frozen, and more time in the exposed centre area (i.e. bolder  
402 fish tended to be more bold across all three variables). Fish showed no change in their response  
403 to the predator strike between replicates (i.e. period\*replicate effects), but showed an overall  
404 increase in activity/ reduction in freezing time across the two replicates (i.e. replicate effects,  
405 Table 4).

406

407 Treatment, i.e., conspecific chemical alarm cue, had a significant positive effect on cortisol  
408 concentration in blood (Fig. 4), but no effect on blood lactate (Table 5). There were no  
409 significant interactions between the treatment effect and the Population, suggesting there is  
410 no difference between populations in the cortisol or lactate responses to a conspecific  
411 chemical cue. Body size and condition also had no effect on cortisol or lactate levels.  
412 Dopaminergic and serotonergic mass ratio was negatively associated with body size, i.e.,  
413 dopaminergic activity was generally lower in larger fish. Similarly, body condition also had a  
414 negative effect on serotonergic mass ratio, i.e., fish in poorer body condition had higher  
415 overall serotonergic activity than higher condition fish. There was no population or  
416 interaction effects dopaminergic and serotonergic mass ratio (Table 5).

417

418 Within both populations, there were not significantly non-zero correlations between  
419 behavioural response variables, and physiological response variables in the cue treatment  
420 group, suggesting that behavioural responses were not directly linked to the measured

421 physiological stress response parameters (for further details see supplementary materials S2,  
422 Table S1 and S2).

423

#### 424 **4. Discussion**

425 Behavioural consistency was found to be high overall across most behavioural variables  
426 measured, but there were some differences in behavioural compositions of each population.  
427 Repeatability of activity, centre use and freezing time appeared to be strong, relative to the  
428 finding of Bell et al. (2009), who showed in a meta-analysis that approximately 35% of  
429 phenotypic variation in behaviour could be attributed to among-individual differences. We  
430 found boldness in the Grønsund population to be repeatable across several variables (i.e.,  
431 activity, centre use and freezing), although significant repeatability in activity and freezing  
432 was not found in Søvng fish. In Thorlacius et al. (2015), bolder gobies from a newly  
433 established populations showed more repeatable individual differences in behaviour. This is  
434 in contrast with our results, as the Grønsund population seemed to be bolder and more  
435 consistent. Moreover in a study about the relation between personality and metabolism (and  
436 also size) in round gobies from an established population in Guldborgsund (first recorded in  
437 2009), latency to resume movement was found to be repeatable (Behrens et al., 2020).

438

439 In contrast with our prediction, fish from Søvng appeared more affected by (i.e., decreased  
440 their activity more in response to) the predator strike than the fish from Grønsund. In  
441 Thorlacius and Brodin (2018), a positive correlation was found between activity and boldness  
442 in round gobies. In that study, the experimental design was the same as ours. Thus, fish from  
443 Grønsund might be considered bolder, different from our original prediction. In another study  
444 on round gobies, Groen et al. (2012) did not find any significant difference in boldness  
445 between old and new populations. Notably, our newer population was older (5 years post-  
446 establishment) than comparable studies (i.e. 2-3 year; Myles-Gonzalez et al. 2015; Groen et  
447 al. 2012), so that difference observed in those studies may have been lost by the time of  
448 sampling. Traits linked to dispersal like boldness may be lost over time due to differences in  
449 density between populations: lower densities lead to selection of traits that increase  
450 reproductive rate rather than dispersal tendency (Duckworth, 2008). The type of behavioural  
451 variables being considered may also be a factor leading to our contrasting results, for  
452 example only Groen et al. (2012) measured boldness as activity after a stressful event, while  
453 Myles-Gonzalez et al. (2015) measured it as the latency to emerge from a shelter.

454

455 Contrary to expectations, there was no difference between populations in their physiological  
456 stress responses. The fish did show a stress response after exposure to the cue, as demonstrated  
457 by the blood cortisol concentration increase, however, there was no effect of the cue on lactate.  
458 No significant changes in lactate levels in blood suggests that no anaerobically fueled activity  
459 was triggered. While high swimming activity may occur as part of the behavioural response to  
460 stress in fish (Wells & Pankhurst, 1999), in this case, the response was primarily freezing with  
461 limited swimming activity. This may be a characteristic of this species and/or population, or  
462 potentially a result of our specific experimental setup.

463

464 We predicted an effect of condition on behavioural responses to predator cues, but only a small  
465 effect on centre use was found, where fish in better condition were less responsive to the  
466 predator strike. This result is in contrast with Moran et al. (2021), who showed a negative effect  
467 of nutritional condition on boldness, where fish subject to poor feeding treatments tended  
468 engage in more risk-taking behaviour than well fed fish. However, in the present study, no  
469 manipulation of the feeding regime was performed, the two populations did not show  
470 difference in condition, and variation within populations was low. Moreover, the long time  
471 spent in the holding facility could have affected the general condition of the fish, that in nature  
472 is likely to be more variable, thus the only effect we detected was small and on only one of the  
473 behaviour variables (centre use). Total length, instead, showed significant effects on every  
474 behaviour variable. Total length showed a very strong negative effect on the activity in a novel  
475 environment and a minor non-significant negative effect on the activity in a predator response  
476 context: smaller fish were more active overall, but they showed greater reduction in activity  
477 after the strike. Movement into refuge areas is a typical response of animals to a predator  
478 presence and moving into the corner areas of the arenas may reflect this behaviour given there  
479 were no sheltered areas in the arena. The negative effect of size on boldness contrasts with the  
480 results from the meta-analysis by Niemelä and Dingemanse's study (2018), as for hormone  
481 levels, Niemelä and Dingemanse found a weak positive correlation between body size and  
482 boldness traits. However, in a species-specific case the results could be different. In fact in  
483 (Behrens et al., 2020), the correlation between size and boldness in round gobies resulted to be  
484 negative. We found a small significant effect of total length was detected on centre use, both  
485 before and after the strike, where larger fish were less responsive to the predator strike, and so  
486 slightly bolder. Finally, we detected a significant effect of total length on freeze time, where  
487 smaller fish spent less time frozen overall, but more after the predator strike. From these two  
488 results, it seems like larger fish are generally less responsive to a predator strike. This could be

489 because they were already less active in the pre-strike period, or maybe because their size  
490 makes them less vulnerable to predation. However, even if the fish closer to the invasion front  
491 are larger, no population effect was detected.

492

493 We found no correlations between physiological and behavioural variables. This is in contrast  
494 with the result from meta-analysis performed by Niemelä and Dingemanse (2018). In that  
495 study, only a weak overall correlation between hormone levels and among-individual  
496 behavioral variation was found (e.g. aggression, boldness, exploration, and activity traits)  
497 across numerous different species. Notably, these are the results of a meta-analysis of many  
498 studies, so the effect for any single species or future study will deviate. Some studies focused  
499 on single species have shown no relationship between stress coping styles (e.g., cortisol level  
500 in the blood) and behavioural responses, e.g. in rainbow trout in acute confinement stress and  
501 novel object response (*Oncorhynchus mykiss*; Gesto, 2019; Thomson et al., 2011) and  
502 seabream exposed to air (*Sparus aurata*, farmed; Höglund et al., 2020). Our data is in line with  
503 these studies and does not support a strong link between behavioral and physiological responses  
504 in this case.

505

506 In summary in invasive round gobies, population differences, body size and body condition  
507 all appear to be factors influencing boldness/risk-taking behaviours, but effects are different  
508 depending on which specific boldness-related behavioural variable is being measured.  
509 Several variables related to boldness/risk-taking behaviour were repeatable across all  
510 experimental fish and within Grønsund fish, but fewer were repeatable within Søvang gobies,  
511 suggesting there may be some differences in behavioural variation within each population.  
512 Exposure to conspecific chemical cues resulted in an increase of cortisol concentration in  
513 blood, but with no differences in responses between the two populations with different  
514 invasion histories. Finally, no correlation between behavioural responses and physiological  
515 responses was detected, and only a small effect of condition on centre use was found.  
516 Furthermore, this study again suggests that among-individual behavioural variation may play  
517 an important role in the invasion process of the round goby, and specifically identifies  
518 boldness in a predator response context as a key behavioural difference between these two  
519 populations. Despite this, physiological variation was not identified as factors directly driving  
520 personality trait variation within or between these two populations. Therefore, valuable future  
521 research may include experimental studies specifically focusing on the role of invasion  
522 processes in driving phenotypic variation along their invasion front (e.g. by analyzing



523 variation across multiple populations), and further in depth analysis of the underlying  
524 mechanisms that produce behavioural variation within populations.

525

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530

## 531 **6. Author Contributions (CRediT taxonomy)**

532 Andrea Galli: Conceptualization, Data curation, Formal Analysis, Investigation,  
533 Methodology, Project administration, Software, Validation, Visualization, Writing – original  
534 draft, Writing – review & editing

535 Jane W. Behrens: Conceptualization, Funding acquisition, Investigation, Methodology,  
536 Project administration, Resources, Supervision, Validation, Writing – review & editing

537 Manuel Gesto: Conceptualization, Data curation, Investigation, Methodology, Resources,  
538 Validation, Writing – review & editing

539 Nicholas P. Moran: Conceptualization, Data curation, Formal Analysis, Funding acquisition,  
540 Investigation, Methodology, Project administration, Resources, Software, Supervision,  
541 Validation, Visualization, Writing – original draft, Writing – review & editing

542

## 543 **7. Data Availability Statement**

544 All data and code used (including data processing, preparation, analysis and presentation) are  
545 available at the Open Science Framework (<https://osf.io/fb8nz/>,  
546 doi:10.17605/OSF.IO/FB8NZ)

547

## 548 **8. Supporting Information**

549 **S1.** Physical composition of experimental populations

550 - Fig. S1, condition, length and weight comparisons between populations.

551 **S2.** Correlation Matrices

552 - Table S1, correlation matrix, Søvvang.

553 - Table S2, correlation matrix, Grønsund

554

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**Table 1.** Behavioural variables recorded from predator response experiments, including the period they were collected in, and the distributions/transformations used in analysis.

	Phase	Distribution	Description
<i>Activity</i> (mm)	<i>pre-strike/</i>	Gaussian	Total distance moved by the fish during each 5 min period.
	<i>post-strike</i>	( <i>sqrt(x)</i> transformed)	Greater movement can represent increased boldness/ risk-taking.
<i>Centre use</i> (s)	<i>pre-strike/</i>	Poisson	The number of seconds spent more than 5 cm from the arena's edges during each 5 min period. Spending more time in central/exposed areas can represent increased boldness/ risk-taking.
	<i>post-strike</i>		
<i>Freezing time</i> (s)	<i>pre-strike/</i>	Gaussian	Total time frozen during each 5 min period. Freezing event were recorded within ToxTrac as periods of greater than 3 s, where the fish's movement was below 30mm. The minimum speed to be considered mobile was 5 mm/s. More time spend frozen can represent lower boldness/ risk-taking behaviour.
	<i>post-strike</i>	( <i>sqrt(t<sub>max</sub>-x)</i> transformed)	
<i>Latency to first freezing</i> (s)	<i>post-strike</i>	Binomial	Fish that were quick to freeze those with a latency < median latency to first freezing post-strike (i.e. approx 9s). Freezing more quickly in response to a predation cue can represent lower boldness/ risk-taking.
		(1 = freeze < 9 s) 0 = freeze > 9 s)	
<i>Latency to resume movement</i> (s)	<i>post-strike</i>	Binomial	Fish that were quick to resume movement following a freezing event were those with a latency < median latency (i.e. approx. 30s). Based on our assessment of freezing responses from videos, freezing events after 20s appeared unrelated to the strike, so fish that did not freeze within 20s were also scored as 0. Resuming normal behaviour more quickly after an exposure to an acute predator cue is considered to represent higher boldness/risk-taking.
		(1 = recovery > 30 s, 0 = recovery < 30 s)	

**Table 2.** Physiological variables taken from blood and brain samples.

	<b>Distribution</b>	<b>Description</b>
<i>Cortisol (ng/mL)</i>	Gaussian ( $\log_e(x)$ transformed)	The main glucocorticoid hormone in teleost fish, produced and released upon activation of the HPI (hypothalamus-pituitary-interrenal) axis. Plasma cortisol levels are used as primary indicator of stress.
<i>Lactate (mM)</i>	Gaussian ( $\log_e(x)$ transformed)	Product of tissue anaerobic metabolism, used as indicator of behavioral activation during stress.
<i>Dopaminergic mass ratio (%)</i>	Gaussian ( $\log_e(x)$ transformed)	Mass ratio in the telencephalon: DOPAC/dopamine, used as an indirect indicator of dopaminergic neuron firing.
<i>Serotonergic mass ratio (%)</i>	Gaussian ( $\log_e(x)$ transformed)	Mass ratio in the telencephalon: 5HIAA/serotonin, used as an indirect indicator of serotonergic neuron firing.

**Table 3.** Repeatability estimates as a measure of behavioural consistency for each behavioural variable measured, including: among-individual variation in both populations as a proportion of total variance ( $ICC_{Raw}$ ); the proportion of among-individual variance excluding population-level variation ( $ICC_{Adjusted}$ ); and estimates within the older ( $ICC_{Grønsund}$ ) and more recently established ( $ICC_{Søvang}$ ) populations. Values in bold represent significantly non-zero coefficients of repeatability.

Variable	Phase	$ICC_{Raw}$	$ICC_{Adjusted}$	$ICC_{Grønsund}$	$ICC_{Søvang}$
<i>Activity (mm)</i>	<i>pre-strike</i>	<b>0.47 [0.25, 0.61]</b> ( <b>p &lt; 0.001</b> )	<b>0.42 [0.17, 0.63]</b> ( <b>p &lt; 0.001</b> )	<b>0.58 [0.35, 0.75]</b> ( <b>p &lt; 0.001</b> )	0.28 [0, 0.59] (p = 0.085)
	<i>post-strike</i>	<b>0.39 [0.2, 0.58]</b> ( <b>p &lt; 0.001</b> )	<b>0.26 [0.06, 0.51]</b> ( <b>p = 0.005</b> )	<b>0.42 [0.11, 0.62]</b> ( <b>p = 0.008</b> )	0.2 [0, 0.47] (p = 0.143)
<i>Centre use (s)</i>	<i>pre-strike</i>	<b>0.54 [0.29, 0.72]</b> ( <b>p &lt; 0.001</b> )	<b>0.54 [0.33, 0.68]</b> ( <b>p &lt; 0.001</b> )	<b>0.36 [0, 0.59]</b> ( <b>p = 0.032</b> )	<b>0.67 [0.27, 0.81]</b> ( <b>p &lt; 0.001</b> )
	<i>post-strike</i>	<b>0.59 [0.43, 0.7]</b> ( <b>p &lt; 0.001</b> )	<b>0.57 [0.36, 0.77]</b> ( <b>p &lt; 0.001</b> )	<b>0.34 [0.06, 0.6]</b> ( <b>p = 0.035</b> )	<b>0.74 [0.47, 0.86]</b> ( <b>p &lt; 0.001</b> )
<i>Freezing time (s)</i>	<i>pre-strike</i>	<b>0.39 [0.2, 0.55]</b> ( <b>p = 0.001</b> )	<b>0.34 [0.14, 0.49]</b> ( <b>p = 0.002</b> )	<b>0.48 [0.17, 0.69]</b> ( <b>p = 0.003</b> )	0.21 [0, 0.5] (p = 0.154)
	<i>post-strike</i>	<b>0.44 [0.21, 0.61]</b> ( <b>p &lt; 0.001</b> )	<b>0.3 [0.1, 0.5]</b> ( <b>p = 0.001</b> )	<b>0.47 [0.15, 0.67]</b> ( <b>p = 0.003</b> )	0.23 [0, 0.53] (p = 0.114)
<i>Latency to first freezing (s)</i>	<i>post-strike</i>	0 [0, 0.17] (p = 0.5)	0 [0, 0.09] (p = 1)	0 [0, 0.22] (p = 1)	0.01 [0, 0.26] (p = 0.456)
<i>Latency to resume movement (s)</i>	<i>post-strike</i>	0 [0, 0.18] (p = 0.5)	0 [0, 0.17] (p = 0.5)	0 [0, 0.21] (p = 0.5)	0 [0, 0.34] (p = 0.5)

**Table 4.** Effects of SL, condition factor, population and replicate number on behavioural responses. Interactions between these factors and period represent the effect of these factors on the change in behaviour between the *pre-strike* and *post-strike* periods, i.e. effects on their response to the acute predator cue. Positive effects on activity represent increased movement (i.e. higher boldness), positive effects on centre area use represent increased use of the central/exposed area (i.e. higher boldness), and positive effects on freeze time represent reduced time spend frozen (i.e. higher boldness). Fixed effect factors/interactions with a non-zero effect are highlighted in bold. Conditional  $R^2$  ( $R^2_{Con}$ ) represents the proportion of variance explained by random and fixed effects within the model, and Marginal  $R^2$  ( $R^2_{Mar}$ ) represents the proportion of variance explained by the fixed effects.

<i>Model</i>	<i>Estimate [95% CI]</i>	<i>S.E.</i>	<i>df</i>	<i>t/z</i>	<i>P</i>	$R^2_{Con}$	$R^2_{Mar}$
<i>- factors</i>							
Activity <sub>(sqrt(x), gaussian lmer model)</sub>						0.575	0.157
- <i>intercept</i>	75.62 [65.07, 86.15]	5.46	76.84	13.85	< 0.001***		
- <i>period</i>	-0.38 [-8.75, 7.99]	4.32	180.18	-0.09	0.93		
- <b>length<sub>Z-scaled</sub></b>	<b>-13.43 [-21.01, -5.84]</b>	<b>3.93</b>	<b>85.31</b>	<b>-3.41</b>	<b>&lt; 0.001***</b>		
- <i>condition<sub>Z-scaled</sub></i>	-1.36 [-7.91, 5.2]	3.4	87.61	-0.4	0.69		
- <i>population<sub>Søvang</sub></i>	-2.78 [-17.74, 12.17]	7.75	85.53	-0.36	0.721		
- <b>replicate<sub>Z-scaled</sub></b>	<b>5.31 [1.28, 9.3]</b>	<b>2.07</b>	<b>182.35</b>	<b>2.57</b>	<b>0.011*</b>		
- <b>period*length<sub>Z-scaled</sub></b>	<b>10.12 [3.7, 16.54]</b>	<b>3.31</b>	<b>180.18</b>	<b>3.06</b>	<b>0.003**</b>		
- <i>period*condition<sub>Z-scaled</sub></i>	2.62 [-3.02, 8.26]	2.91	180.18	0.9	0.368		
- <b>period*population<sub>Søvang</sub></b>	<b>-15.16 [-27.97, -2.36]</b>	<b>6.6</b>	<b>180.18</b>	<b>-2.3</b>	<b>0.023*</b>		
- <i>period*replicate<sub>Z-scaled</sub></i>	-0.19 [-5.82, 5.45]	2.91	180.18	-0.06	0.949		
Centre use <sub>(count, poisson glmer model)</sub>						0.984	0.039
- <i>intercept</i>	3.51 [2.9, 4.12]	0.31	n/a	11.42	< 0.001***		
- <b>period</b>	<b>-0.37 [-0.42, -0.31]</b>	<b>0.03</b>	<b>n/a</b>	<b>-13.01</b>	<b>&lt; 0.001***</b>		
- <i>length<sub>Z-scaled</sub></i>	0.03 [-0.42, 0.49]	0.23	n/a	0.14	0.887		
- <i>condition<sub>Z-scaled</sub></i>	-0.12 [-0.52, 0.27]	0.2	n/a	-0.63	0.527		
- <i>population<sub>Søvang</sub></i>	-0.59 [-1.51, 0.3]	0.45	n/a	-1.3	0.192		
- <i>replicate<sub>Z-scaled</sub></i>	-0.02 [-0.05, 0]	0.01	n/a	-1.88	0.06		
- <b>period*length<sub>Z-scaled</sub></b>	<b>0.05 [0.01, 0.09]</b>	<b>0.02</b>	<b>n/a</b>	<b>2.49</b>	<b>0.013*</b>		
- <b>period*condition<sub>Z-scaled</sub></b>	<b>0.07 [0.03, 0.11]</b>	<b>0.02</b>	<b>n/a</b>	<b>3.61</b>	<b>&lt; 0.001***</b>		
- <i>period*population<sub>Søvang</sub></i>	0.03 [-0.05, 0.12]	0.04	n/a	0.71	0.48		
- <i>period*replicate<sub>Z-scaled</sub></i>	0 [-0.04, 0.04]	0.02	n/a	-0.06	0.956		
Freeze time <sub>(sqrt(tmax-x), gaussian lmer model)</sub>						0.578	0.159
- <i>intercept</i>	9.95 [8.52, 11.38]	0.74	79.24	13.43	< 0.001		
- <i>period</i>	-0.08 [-1.21, 1.06]	0.59	181.23	-0.13	0.897		
- <b>length<sub>Z-scaled</sub></b>	<b>-1.56 [-2.6, -0.53]</b>	<b>0.54</b>	<b>85.57</b>	<b>-2.91</b>	<b>0.005**</b>		
- <i>condition<sub>Z-scaled</sub></i>	-0.3 [-1.19, 0.6]	0.46	87.85	-0.64	0.522		
- <i>population<sub>Søvang</sub></i>	-0.72 [-2.77, 1.32]	1.06	85.78	-0.68	0.496		
- <b>replicate<sub>Z-scaled</sub></b>	<b>0.98 [0.43, 1.52]</b>	<b>0.28</b>	<b>183.37</b>	<b>3.49</b>	<b>&lt; 0.001***</b>		
- <b>period*length<sub>Z-scaled</sub></b>	<b>1.02 [0.15, 1.89]</b>	<b>0.45</b>	<b>181.23</b>	<b>2.27</b>	<b>0.025*</b>		
- <i>period*condition<sub>Z-scaled</sub></i>	0.57 [-0.19, 1.34]	0.4	181.23	1.45	0.148		
- <i>period*population<sub>Søvang</sub></i>	-1.67 [-3.41, 0.07]	0.9	181.23	-1.86	0.065		
- <i>period*replicate<sub>Z-scaled</sub></i>	-0.16 [-0.92, 0.61]	0.39	181.23	-0.4	0.691		

**Table 5.** Effects of fish length, condition factor, population and alarm cue treatment on physiological variables. Interactions between population and treatment are used to test if the response to an alarm cue (i.e. *sham control* versus *cue* groups) differs between populations (i.e. Grønsund and Søvang). Fixed effect factors/interactions with a non-zero effect are highlighted in bold. Conditional  $R^2$  ( $R^2_{\text{Con}}$ ) represents the proportion of variance explained by random and fixed effects within the model, and Marginal  $R^2$  ( $R^2_{\text{Mar}}$ ) represents the proportion of variance explained by the fixed effects.

<i>Model</i>							$R^2_{\text{Con}}$	$R^2_{\text{Mar}}$
- factors	<i>Estimate [95% CI]</i>	<i>S.E.</i>	<i>df</i>	<i>t/z</i>	<i>P</i>			
<b>Cortisol</b> <sub>(log(x), gaussian lmer model)</sub>							0.223	0.132
- <i>intercept</i>	2.9 [2.4, 3.4]	0.27	5.95	10.56	< 0.001***			
- <b>treatment</b>	<b>0.51 [0.06, 0.95]</b>	<b>0.23</b>	<b>43.83</b>	<b>2.16</b>	<b>0.036*</b>			
- <i>length</i> <sub>Z-scaled</sub>	0.01 [-0.26, 0.27]	0.14	45.48	0.05	0.958			
- <i>condition</i> <sub>Z-scaled</sub>	-0.07 [-0.31, 0.16]	0.12	46.74	-0.58	0.562			
- <i>exposure time</i>	-0.2 [-0.66, 0.27]	0.24	44.31	-0.81	0.42			
- <i>population</i> <sub>Søvang</sub>	-0.34 [-1.01, 0.35]	0.37	4.8	-0.92	0.403			
- <i>treatment*population</i> <sub>Søvang</sub>	-0.25 [-0.91, 0.43]	0.35	43.95	-0.71	0.481			
<b>Lactate</b> <sub>(log(x), gaussian lmer model)</sub>							0.085	0.081
- <i>intercept</i>	-1.82 [-2.08, -1.57]	0.14	14.54	-13.18	< 0.001***			
- <i>treatment</i>	-0.07 [-0.35, 0.21]	0.15	46.05	-0.46	0.644			
- <i>length</i> <sub>Z-scaled</sub>	0.09 [-0.07, 0.26]	0.09	46.8	1.05	0.3			
- <i>condition</i> <sub>Z-scaled</sub>	-0.07 [-0.22, 0.08]	0.08	47.99	-0.9	0.372			
- <i>exposure time</i>	0.27 [-0.03, 0.56]	0.16	46.58	1.7	0.095			
- <i>population</i> <sub>Søvang</sub>	-0.15 [-0.47, 0.18]	0.18	46.26	-0.83	0.41			
- <i>treatment*population</i> <sub>Søvang</sub>	-0.01 [-0.43, 0.41]	0.23	46.12	-0.03	0.974			
<b>Dop. mass ratio</b> <sub>(log(x), gaussian lmer model)</sub>							0.274	0.155
- <i>intercept</i>	1.39 [1.09, 1.7]	0.17	5.75	8.38	< 0.001***			
- <i>treatment</i>	-0.05 [-0.3, 0.2]	0.13	43.97	-0.41	0.681			
- <b>length</b> <sub>Z-scaled</sub>	<b>-0.21 [-0.36, -0.06]</b>	<b>0.08</b>	<b>45.32</b>	<b>-2.73</b>	<b>0.009**</b>			
- <i>condition</i> <sub>Z-scaled</sub>	-0.06 [-0.19, 0.08]	0.07	46.4	-0.84	0.402			
- <i>exposure time</i>	-0.22 [-0.48, 0.04]	0.14	44.36	-1.63	0.109			
- <i>population</i> <sub>Søvang</sub>	0.29 [-0.12, 0.71]	0.22	4.79	1.3	0.254			
- <i>treatment*population</i> <sub>Søvang</sub>	-0.06 [-0.44, 0.31]	0.2	44.06	-0.3	0.766			
<b>Ser. mass ratio</b> <sub>(log(x), gaussian lmer model)</sub>							0.328	0.302
- <i>intercept</i>	2.95 [2.84, 3.06]	0.06	7.37	47.98	< 0.001***			
- <i>treatment</i>	0.03 [-0.09, 0.15]	0.06	43.96	0.48	0.632			
- <b>length</b> <sub>Z-scaled</sub>	<b>-0.12 [-0.19, -0.05]</b>	<b>0.04</b>	<b>46.26</b>	<b>-3.31</b>	<b>0.002**</b>			
- <b>condition</b> <sub>Z-scaled</sub>	<b>-0.09 [-0.15, -0.03]</b>	<b>0.03</b>	<b>47.67</b>	<b>-2.83</b>	<b>0.007**</b>			
- <i>exposure time</i>	0.03 [-0.09, 0.15]	0.06	44.65	0.52	0.605			
- <i>population</i> <sub>Søvang</sub>	-0.03 [-0.18, 0.12]	0.08	5.46	-0.4	0.708			
- <i>treatment*population</i> <sub>Søvang</sub>	-0.04 [-0.21, 0.14]	0.09	44.12	-0.41	0.687			

780 **Figure Legends**

781 *Figure 1.* Study sites in on the eastern coast of Denmark, and inset, within the broader south-  
782 east Baltic Sea region. Round gobies were first recorded in the areas around Grønsund (pink)  
783 and Søvng (green) in approximately 2011 and 2016, respectively, 10 and 5 years prior to  
784 sampling. Grønsund is located directly north of Guldborgsund, the first area of inner Danish  
785 waters invaded by round gobies in 2009. Søvng is located just south of Copenhagen, nearer  
786 to the DTU Lyngby laboratory in northern Copenhagen. At present, DTU Lyngby is  
787 approximately level with the northern edge of the round goby's invasion front up the eastern  
788 coast of Denmark.

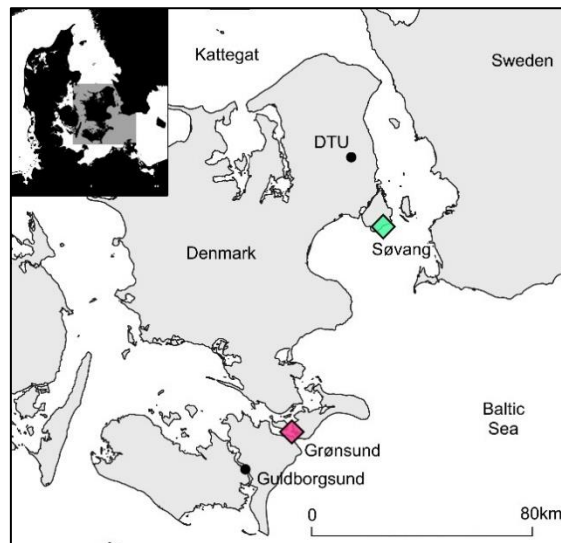
789  
790 *Figure 2.* Behavioural arenas (a) aerial view, and (b) front view. In (a), the grey area  
791 represents the 5cm edge zone used to measure centre area use behaviour, and the parallel  
792 lines across the arenas represent the clear acrylic tube used to suspend the bolt above the  
793 centre of the arena. The fishing line runs through the length of the tube and can be released  
794 and retracted by the observer. The bolt-release mechanism sits on top of a frame used to hold  
795 the polystyrene walls around the arena (dotted lines), and to hold the camera in position, so  
796 that the bolt drops approximately 80cm to break the water surface before retraction.

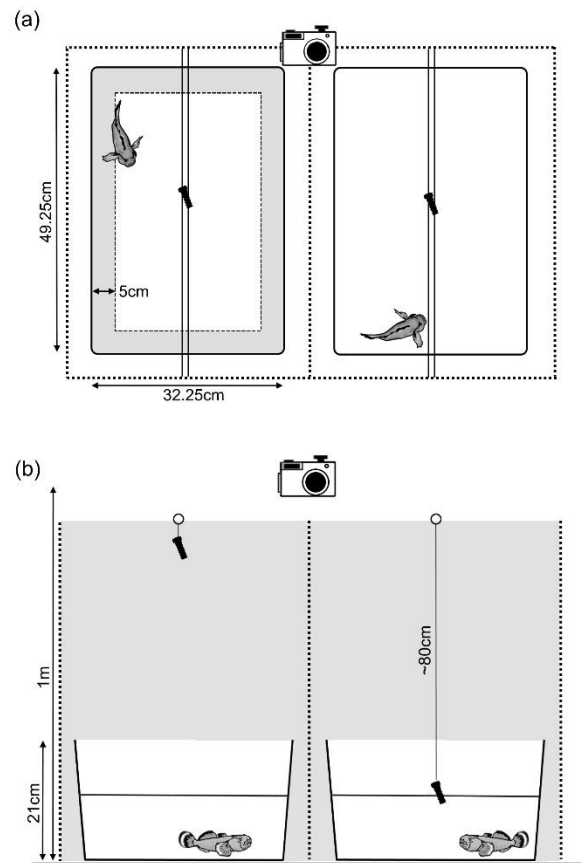
797  
798 *Figure 3.* Activity response in Experiment 1, associated with (a) population, and (b) SL. In  
799 (a) fish from the newer site Søvng (est. 2016) appear to reduce their activity levels following  
800 the predator cue more than Grønsund (est. 2011) fish. In (b) smaller fish were more active  
801 overall but reduced their activity levels more relative to larger fish following the predator  
802 cue. In boxplots horizontal bars, boxes and vertical lines represent the median, interquartile  
803 range, non-outlier range, respectively.

804  
805 *Figure 4.* Blood cortisol response to a chemical alarm cue. The dotted line represents the  
806 baseline measurements for cortisol in the *baseline control* treatment group. Concentrations  
807 are significantly higher in the *cue* treatment group, but there is no difference between  
808 populations. In boxplots horizontal bars, boxes and vertical lines represent the median,  
809 interquartile range, non-outlier range, respectively.

810

Figure 1



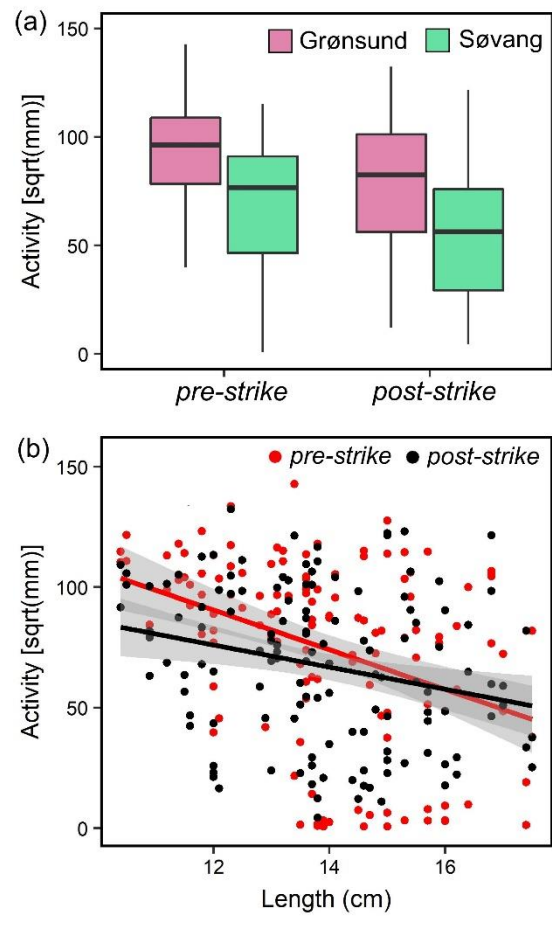




813

Figure 3

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815

Figure 4

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