1 Visual and olfactory cues of predation affect body and brain growth

2 in the guppy

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10 Abstract

- Phenotypic plasticity requires animals to acquire reliable environmental information.
 When multiple sources of information agree, cues should be perceived as reliable and induce a relatively strong response. Conversely, where stimuli conflict, animals must weigh the accuracy of the sources of information and responses should be reduced.
 Availability of reliable information is often considered a limitation on plasticity, yet how animals integrate seemingly contradictory or incomplete information remains enigmatic, as empirical tests are scarce.
- We tested how incomplete information determines phenotypic plasticity by
 simulating predation risk during early ontogeny of guppies (*Poecilia reticulata*). We
 exposed guppy fry to a combination of visual and/or olfactory cues of the predatory
 pike cichlid (*Crenicichla alta*), and monitored growth of the body and brain. After five
 weeks of exposure, guppies were returned to common no-risk conditions and their
 activity rates were monitored for four weeks post-treatment.
- Visual predator exposure more strongly affected development; reducing body size of
 adult males and increasing brain size in females. However, there was little evidence
 for the hypothesised additive effect, with the combined treatment not inducing a
 larger effect than when only receiving olfactory or visual treatments.
- 28 5. While there was consistent individual variation in activity rates, this was unaffected
 29 by developmental risk and uncorrelated with the growth parameters.
- Our results demonstrate the differential reliability of cues during development. Visual
 exposure to a predator was a highly reliable environmental cue, while environmental
 certainty was unaffected by combined stimuli.

Key words: Behavioural plasticity, developmental plasticity, animal personality, pace of life
 syndrome, U-model,

35 Introduction

36 Environmental conditions are constantly changing. While many factors change predictably 37 (e.g. seasonal change), other factors are much less predictable, such as movement and 38 density of predators. Accordingly, animals survey the environment to gather information 39 which informs the development of various phenotypic traits, including behaviour (Stein et 40 al., 2018; Urszán Tamás et al., 2018), life-history (Torres-Dowdall et al., 2012), and 41 morphology (Agrawal, 2001). Information often comes from multiple sources which may 42 conflict temporally, for example when parental and individual experience differs (Salinas 43 and Munch, 2012; Stein et al., 2018) or if information is sampled from different cues (e.g. 44 visual, olfactory, auditory, and social cues) (Munoz and Blumstein, 2012). Animals must 45 therefore gauge the accuracy of acquired information, as mismatches between the 46 environmental conditions and expressed phenotype can have high fitness costs (Nussey et 47 al., 2005).

48 Environmental information processing can be conceptualised as a Bayesian updating 49 process (Stamps and Frankenhuis, 2016). Animals begin life with a prior 'belief' of the state 50 of the environment, derived from their ancestors (genetic and transgenerational plasticity), 51 which is updated in response to information from the current environmental state -52 represented by a likelihood distribution (Stamps and Bell, 2020; Stamps and Krishnan, 53 2014). Where multiple stimuli indicate the same conditions, this could be represented as 54 high certainty in the environmental state, and thus induce a stronger phenotypic response 55 relative to a solitary cue. Conversely, when stimuli conflict, animals must weigh the relative 56 reliability that the stimulus (or absence of the stimulus) conveys to inform the plastic 57 response (Stamps and Bell, 2020). For example, a recent study found that direct visual cues 58 of a predator had a greater effect on behaviour than indirect alarm and mobbing calls in 59 adult black-capped chickadees (Poecile atricapillus) - indicating visual cues were perceived 60 as more reliable (Arteaga-Torres et al., 2020). Such evaluation of the relative certainty of 61 stimuli should become even more important over developmental timeframes and in traits 62 where plasticity is non-reversible, as responses are predictive of future adult environmental 63 state.

Developmental studies of predation usually manipulate one stimulus – often olfactory
 cues in aquatic systems (Agrawal, 2001; Brönmark and Pettersson, 1994; Urszán Tamás et

66 al., 2018). This stimulus is assumed to provide information about the state of the 67 environment, to estimate the density or relative risk of predators (Stamps and Bell, 2020). 68 By extension, the absence of cues is implicitly assumed to convey low density (or absence) 69 of predators (e.g. Ghalambor et al., 2015), which is often used as the control. Such olfactory 70 cues can also be multifaceted, conveying different pieces of information. In crucian carp 71 (*Carassius carassius*), olfactory cues of a predator lead to a plastic response of increased 72 body depth as a defence against gape-limited predators (Brönmark and Pettersson, 1994), 73 though skin extract cues isolated from tissue extract cues failed to replicate the effect 74 (Stabell et al., 2010). Alternative study designs have manually chased fish to simulate risk 75 (Edenbrow and Croft, 2013), added visual cues (Reddon et al., 2018; Stein et al., 2018), or 76 replicated the social environment to resemble contrasting predation regimes (Rodd et al., 77 1997). In natural systems, while these cues are not independent (e.g. olfactory cues will 78 positively correlate with visual encounter rate), different life-stages or individuals may 79 occupy different areas and not always receive complete information. Visual encounter rates 80 or olfactory cue concentrations alone may therefore be insufficient to predict the wider environment. Multiple cues could add greater certainty to the perceived environmental 81 82 state, while also providing subtly different information. Studies which contrast multiple 83 sources of environmental information have been well considered in the immediate 84 environment (Munoz and Blumstein, 2012), though are relatively lacking in developmental 85 plasticity.

86 In this experiment, we aimed to quantify the effect of conflicting or incomplete 87 information on development, both from different sensory modalities and a temporal change 88 in environmental state. We make use of the guppy system, where there are known effects 89 of high predation – commonly defined as the presence of the Trinidadian pike cichlid 90 (Crenicichla sp.) (Magurran, 2005). Pike cichlids target larger guppies (Johansson et al., 91 2004) which adds a strong selection pressure on life-history traits, increasing growth rates 92 while decreasing size at birth, size at maturity and adult size (Reznick and Endler, 1982; Reznick et al., 1990). Predation risk also has broad behavioural effects, increasing gregarious 93 94 behaviour (Seghers, 1974), and affecting social structure (Herbert-Read et al., 2017). Fish 95 from high predation environments are slower to habituate to a novel stimulus (Brown et al., 96 2013) and make decisions slower than low predation fish (Burns and Rodd, 2008).

Accordingly, there are associated changes in brain morphology (though reported effects are
variable) (Kotrschal et al., 2017; Mitchell et al., 2020b; Reddon et al., 2018). Artificial
selection on brain size has revealed that large-brained fish out-perform small-brained fish in
various cognitive tasks (Buechel et al., 2018; Kotrschal et al., 2013), are better able to assess
risk (van der Bijl et al., 2015) and survive longer when faced with predation (Kotrschal et al.,
2015).

103 Beyond the pronounced geographic variation in predation regimes (Endler, 1978), there 104 is also considerable temporal variation (Deacon et al., 2018), and dispersal of individuals 105 across predator barriers (Crispo et al., 2006). Accordingly, guppies incorporate 106 environmental information during early development into their phenotypes (Handelsman et 107 al., 2013; Torres-Dowdall et al., 2012), though effects often counter the evolved response 108 (Ghalambor et al., 2015). A previous study to evaluate the effect of predation on 109 development of brain anatomy found an increase in relative brain size - consistent with the 110 ecological comparisons of the same study – yet this difference was limited to males (Reddon 111 et al., 2018). Such alterations in relative brain size can result from broader selection or 112 alterations in life-history – either growth rates or adult size (Rogell et al., 2020). These brain anatomy and life-history traits are energetically costly, with predicted associations with 113 114 activity and other behaviours which underly acquisition of resources and energy 115 expenditure (Careau et al., 2008).

116 We exposed guppies to two different sensory cues of the predatory pike cichlid (C. alta), 117 so that cues were either combined, or in isolation. We quantified growth in body and brain 118 size throughout the experiment. Fish were then returned to risk free conditions, causing a 119 temporal change in environmental state. We measured activity rates after fish were 120 returned to risk free conditions, to assess if behavioural differences were retained. These 121 data allowed us to test whether olfactory and visual predation cues additively affected 122 developmental plasticity. We expected predator exposed fish to have decreased growth 123 rates and adult size (Handelsman et al., 2013), an increase in brain size (Reddon et al., 2018) 124 and a decrease in activity rates to reduce encounter rates with predators (Stamps, 2007). 125 More specifically, we predicted the combination of visual and olfactory risk to provide 126 greater environmental certainty and therefore elicit a stronger phenotypic response. In the

127 behavioural assays, where change is likely reversible, we expected treatment groups to

128 converge following the completion of the treatment phase of the experiment.

129

130 Methods

131 Experimental design

Wild-type Trinidadian guppies (*Poecilia reticulata*) were sampled from a stock bred population at Stockholm University. The parents of the focal animals were housed in pairs in nanotanks (12×20×13cm; 4L of water). Each of these tanks contained gravel substrate and Java moss (*Taxiphyllum barbieri*) to provide environmental enrichment, snails (*Planorbis sp.*), and a constant stream of air. Plastic mesh was provided on one side of the tank to provide refuge for fry. Fish were fed twice daily with either flake or live brine *nauplii*.

We checked for fry daily, and when found recorded the date of birth and moved fry to a holding nanotank. Once enough fry were available, fry were photographed (see below) and split into the four treatments. No more than one fry from a mother was allocated to each treatment combination to avoid pseudoreplication.

142 **Developmental treatment**

143 Fish were exposed to the cues of predation for five weeks. Experimental aquaria were 144 designed to expose fry to visual and olfactory stimuli in a full-factorial design (see Fig. 1 for 145 schematic). Fry were held in 16 nanotanks within two pairs of larger tanks (55L). One of the 146 tanks housed a pike cichlid (C. alta) that was fed one freshly culled guppy per day to provide 147 predator chemical cues and possibly information on its diet, and the other was vacant (Fig. 148 1a). The predator was provided shelter in the middle of the tank (shelter also present in the 149 vacant tank). The four nanotanks within the predator tank had visual cues from the 150 predator, while the four nanotanks in the vacant tank were not exposed to visual cues of 151 predation. The two main tanks were connected via two tubes and a pump circulating water 152 between the two tanks, to allow olfactory cues to flow into the vacant tank. Nanotanks from 153 the olfactory treatment had a 1cm hole drilled on the side to allow water to pass, and a fine 154 mesh was glued over the hole to keep fry in. The control nanotanks for olfactory cues were 155 left undrilled. Nanotanks contained between four and seven fish, snails, Java moss, and 156 gravel as substrate.

Fry were reared in this setup for five weeks (see Fig. 1c for timeline), where they were fed daily with hydrated flake food, supplemented with live brine *nauplii* every second day. After five weeks, fish were returned to standard housing conditions and held in groups of 3-4. Fish were sexed based on morphological traits and colouration and kept in single sex groups. A total of four replicates were run from the two setups. 95 fish were used and distributed in the four treatments (Control-Control n = 23; Visual-Control n = 24; Control-Olfactory n =24; Visual-Olfactory n =24).



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Figure 1. Schematic of the experimental setup (a), with treatments indicated in the
nanotanks. The predator was allowed to move freely in its tank. Water was pumped
between main tanks (blue arrows) to circulate predator olfactory cues, and holes were
drilled in the side of olfactory treatment nanotanks. We quantified growth by measuring
body length, abdomen area, and brain area (b). Fish were exposed to the treatment for 5
weeks, we photographed fish nine times across 14-17 weeks to obtain measurements, and
recorded activity three times per burst at weeks 5, 7 and 9 (c).

172 Body and brain growth

173 To quantify growth rates, photos were taken weekly with a Nikon DSLR camera equipped

174 with a Tamron 90mm macro lens. Fish were lit from underneath and photographed from

- above to enhance contrast of the brain. A picture was taken prior to putting the fry in the
- setup and then weekly during the manipulation phase (Fig. 1c). Photos were then taken
- 177 biweekly during the behavioural phase, and a final photo was taken at the end of the
- 178 experiment (14-17wks). From each photo, we measured standard length, abdomen area
- 179 (area of the body from the pectoral fins to the tail fin) and brain area with ImageJ (Schneider
- 180 et al., 2012)(Fig. 1b). Abdomen area aimed to capture variation in body condition and

181 growth not captured by the length measurements. Individual guppies were identified by

182 manually their characteristic melanic spots on their brain and body (see Castillo et al., 2018).

183 Measurements and identification were all performed by a single experimenter (J.L.)

184 Activity assays

185 After conclusion of the treatment phase (week 5), fish were removed from the set-up to 186 record their activity. To do so, fish were placed individually in small arenas (11×11×9cm) 187 with 200mL of water with Java moss for enrichment and left overnight. The activity assay 188 started at 2pm the next day; Java moss was removed, and the arenas were placed onto a 189 filming stage, backlit with infrared light and left to acclimate for 15 minutes. After 190 acclimation, fish were recorded with a USB camera (ELP-USB 100w05MT-SFV) for 30 191 minutes. At the end of the trial, fish were returned to the holding rack still in their activity 192 arena. Moss was placed in the arena again and fish were fed with flake. Videos were recorded for three consecutive days, after which the fish were put back into their holding 193 194 nanotanks in small groups (3-4 fish). Recordings were filmed at 10fps (using OBS Studio) and 195 videos were later tracked with EthoVision XT 10.

This protocol of three activity measures was then repeated for weeks 7 and 9 (i.e. three "bursts" of trials, Fig. 1c). Observations of behaviour taken closely together in time can inflate estimations of individual variation (Mitchell et al., 2020a), so this burst design allows us to better account for potential lack of independence. In the first replicate, 10 fish jumped out of the setup overnight, and subsequently transparent covers were placed on top of the boxes at all times.

Activity rates were measured as the total distance moved for the trial. We analysed a total of N=85 individuals that were recorded during three bursts of observation ($N_{ID:Burst}$ = 204 255) with three observations per burst (N_{obs} = 765).

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206 Statistical analysis

207 Growth data

To assess the effect of predation treatments on growth rates, we fit the data to a Unified Richards growth curve. This equation is a re-parameterisation of the Richards-Bertalanffy

model which removes the mathematical dependencies of the different growth parameters
(Tjørve and Tjørve, 2017). This facilitates comparability of the estimated coefficients
between groups, and correlations of parameters to be assessed among individuals. The
change in size (*S*) as a function of time (*t*) is given by the equation:

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$$S = S_{\infty} \left(1 + \left(\left(\frac{S_0}{S_{\infty}} \right)^{-3} - 1 \right) * exp \left(-4^{\frac{4}{3}}k * t \right) \right)^{\frac{1}{3}}$$
(1)

where S_0 is the size at birth and S_∞ is the asymptotic adult size. The growth coefficient is given by k, and in a unified-model can be interpreted as the maximum relative growth rate at the point of inflexion (Thorley and Clutton-Brock, 2019; Tjørve and Tjørve, 2017). This growth curve is selected from Tjørve and Tjørve (2017) and has a shape parameter set as a constant which moves the point of inflexion higher than the standard von Bertalanffy growth equation – a characteristic which offered a much better fit to the data at hand.

221 The growth model was specified for body length, abdomen area and brain area, with 222 time being the age of the fish in weeks (t). This was fit as a non-linear multivariate mixed 223 model in the Bayesian package brms 3.6.1 (Bürkner, 2017). The S_{∞} and k parameters were 224 fit with the fixed predictors of sex, olfactory treatment, visual treatment and all interactions. 225 As S_0 corresponds to size before any exposure to the treatment, this was fit with the fixed 226 predictor of sex. To evaluate individual variance, identity was fit as a random effect for all 227 three parameters. Correlations between growth parameters were assessed with an 228 unstructured correlation matrix that evaluates all correlations of the three coefficients for 229 the three traits (i.e. a 9×9 matrix).

230 As all three coefficients are bound by 0 (negative size or growth is impossible), 231 coefficients were log-linked to constrain the model to possible values, and post hoc 232 diagnostic plots confirmed this log-normal distribution. Priors were diffuse and directly set 233 on the group means by fixing the global intercept of 0 (see supplementary material for 234 priors and parameterisation). Posteriors were then reorganised to give treatment group 235 comparisons. Residual error was assumed to be normal, though there was a clear pattern of 236 variance expanding. Accordingly, we fit a model for the residual dispersions, with the fixed 237 effect of age.

238 Behavioural data

239 Activity rates were log-transformed to achieve normality, then standardised to a mean of 240 0 and standard deviation of 1. This was fit to the predictors of sex, olfactory treatment, 241 visual treatment, burst, and all interactions. Individual identity (ID) was fit as a random intercept as was the interaction of individual identity by burst (ID:Burst). The separate 242 243 random effects at ID and ID:Burst thus account for temporal dependence of observations 244 taken closely together in time. This temporal dependence is thus placed in the denominator of the repeatability equation to yield a "long-term" repeatability; $R = \frac{\sigma_{ID}^2}{\sigma_{ID}^2 + \sigma_{ID}^2 + \sigma_{E}^2}$ (sensu 245 246 Araya-Ajoy et al., 2015).

247 Growth and behaviour correlations

248 Finally, we aimed to assess whether activity rates correlated with the growth parameters. 249 Equivalent growth coefficients were highly correlated across the three size variables (see 250 Results), so analyses were limited to correlations of body length and activity rates. We used 251 a character-state model for the activity measures, which fits three separate intercepts of 252 activity (one for each burst), which allowed us to test whether the hypothesised correlations 253 with growth were temporally dependent. This model aimed to quantify the phenotypic 254 correlations, so we removed treatment effects. These three burst observations are 255 correlated to the three growth parameters (see above) yielding a 6×6 matrix.

256

257 Results

258 Predator cues impact growth

259 The different predator cues impacted male and female growth differently, and appeared 260 limited to the asymptotic size coefficients. In males, the combined effect of the visual and 261 olfactory treatments reduced adult body length, and abdomen area, relative to the control 262 males (no cues). The incomplete information groups (i.e. only visual or only olfactory) were 263 intermediate in body length, but did not differ significantly from control-control or visual-264 olfactory treatments (Table 1a). This makes it difficult to discern whether the effect was 265 additive (as hypothesised), or constrained by statistical power. Brain area was unaffected by 266 predation risk in males.

267 Females exposed to the visual stimulus of the pike cichlid had larger brain areas, while 268 there was no effect of the olfactory treatment on brain area. There were additionally only 269 small effects of the treatments on the body growth parameters; the combined treatments 270 increased body size, while there was an insignificant increase in visual only treatment. The 271 olfactory stimulus when presented alone had no effect on female growth relative to the control. However, they were different from when the visual cue was presented. This 272 273 indicated that the olfactory treatment was perceived as less reliable than the visual 274 treatment.

275 Combined effects present in body size and brain size indicated that for both sexes, there 276 was an increase in relative brain size. In males, this was driven by a decrease in body size of 277 predator exposed fish, while brain size was unaffected. In females, this was due to an 278 increase in absolute brain size, while effects on body size were small.



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Figure 2: Growth curves are shown for body length (a,d), abdomen area (b,e) and brain area
 (c,f), with females shown on top (a-c) and males on the bottom (d-f). Group geometric mean
 trajectories are overlaid in bold, with thin light colours representing predicted individual
 growth.

			В	ody Lengt	h	Ab	domen Aı	rea	Brain Area		
	Parameter		Est	Lower	Upper	Est	Lower	Upper	Est	Lower	Upper
Females	S_0		2.03	2.007	2.052	1.432	1.382	1.482	0.145	0.113	0.176
		Int	-2.856	-2.926	-2.785	-2.161	-2.225	-2.096	-2.577	-2.668	-2.488
	1.	Vis:Cont	-0.004	-0.104	0.093	0.004	-0.088	0.097	0.009	-0.11	0.127
	к	Cont:Olf	-0.035	-0.145	0.07	-0.042	-0.145	0.056	0.087	-0.044	0.215
		Vis:Olf	-0.069	-0.175	0.036	-0.041	-0.181	0.101	-0.094	-0.228	0.04
		Int	2.939	2.907	2.971	3.42	3.35	3.49	1.081	1.022	1.142
	S_{∞}	Vis:Cont	0.035	-0.01	0.079	0.039	-0.062	0.138	0.107	0.024	0.192
		Cont:Olf	-0.03	-0.075	0.018	-0.09	-0.196	0.017	-0.026	-0.111	0.061
		Vis:Olf	0.05	0.003	0.099	0.081	-0.026	0.189	0.11	0.015	0.208
	S_0		1.989	1.968	2.01	1.338	1.29	1.385	0.13	0.101	0.159
		Int	-2.655	-2.736	-2.575	-2.009	-2.082	-1.936	-2.523	-2.628	-2.421
	1-	Vis:Cont	-0.063	-0.168	0.045	-0.073	-0.171	0.027	-0.006	-0.137	0.127
S	к	Cont:Olf	-0.072	-0.17	0.029	-0.077	-0.171	0.017	-0.029	-0.153	0.098
Male		Vis:Olf	0.012	-0.096	0.118	-0.045	-0.144	0.055	0.019	-0.117	0.155
		Int	2.821	2.789	2.854	3.287	3.216	3.357	1.057	0.994	1.125
	C	Vis:Cont	-0.029	-0.074	0.015	-0.109	-0.21	-0.013	-0.003	-0.089	0.083
	\mathfrak{z}_∞	Cont:Olf	-0.023	-0.064	0.019	-0.106	-0.197	-0.016	0.004	-0.08	0.085
		Vis:Olf	-0.051	-0.094	-0.008	-0.143	-0.239	-0.047	-0.042	-0.127	0.043

Table 1: Displayed are the fixed effect predictions with 95% credible intervals of the growth equations. The intercept is the reference control ('Cont') for both the visual ('Vis') and olfactory ('Olf') treatments, with the three other combinations of treatments given in relation to this reference group. Parameters that do not overlap with 0 are presented in bold.

290

291 Correlations between growth parameters

292 There were very strong correlations between equivalent parameters in body length and

abdomen area (Table 2), which was expected as they are physically and mathematically

linked. Correlations of these variables' starting size (S_0) with starting brain size were very

high, though for the growth coefficient (k), and adult size (S_{∞}) estimates were

296 comparatively weak, with correlations much lower than 1. This indicated the potential for

- 297 brain size to respond independently of body size. Finally, while there were few significant
- 298 correlations of S_0 , k and S_∞ parameters, we found that starting brain size correlated
- significantly with growth rates (k) of the body size variables.

	1_	σ = 0	.110																
th	ĸ	0.086	0.136																
-eng	c	-0.0	078	σ=0	0.045														
d y l	\mathbf{J}_{∞}	-0.327	0.195	0.035	0.056														
BC	ç	0.1	.12	0.1	67	σ = 0	.066												
	50	-0.134	0.355	-0.059	0.38	0.056	0.077												
	l,	0.9	47	0.0	009	0.0	68	σ = 0	0.102										
rea	ĸ	0.878	0.986	-0.262	0.296	-0.176	0.311	0.079	0.128										
en A	c	0.0	33	0.8	865	0.1	.09	0.0	98	σ=0	0.096								
ő	\mathfrak{z}^∞	-0.253	0.323	0.737	0.95	-0.14	0.347	-0.204	0.409	0.074	0.121								
Abd	c	0.1	.79	0.1	29	0.9	66	0.	09	0.1	.15	σ = 0	.142						
	\mathcal{S}_0	-0.081	0.418	-0.101	0.356	0.93	0.988	-0.168	0.349	-0.139	0.353	0.119	0.168	_					
	1.	0.595		0.118		0.2	.97	0.5	532	0.1	.13	0.3	68	σ = 0	.073				
a	к	0.19	0.859	-0.286	0.525	-0.121	0.672	0.108	0.833	-0.302	0.535	-0.04	0.711	0.03	0.115				
Are	c	0.1	.67	0.4	174	-0.2	177	0.2	259	0.5	32	-0.2	223	-0.0)95	σ = 0	.075		
rain	\mathfrak{z}_{∞}	-0.147	0.465	0.191	0.708	-0.448	0.107	-0.06	0.554	0.257	0.756	-0.486	0.058	-0.527	0.398	0.055	0.097		
B	C	0.3	84	0.0)59	0.8	805	0.3	66	0.1	.27	0.7	92	0.2	72	0.0	77	σ=0).086
	3 ₀	0.132	0.61	-0.195	0.302	0.686	0.895	0.117	0.593	-0.144	0.381	0.653	0.894	-0.156	0.687	-0.227	0.368	0.071	0.103
		k		S	8	S	0	ŀ	k	S	8	S	0	k		S	×		S_0
		Body Length					Abdomen Area				Brain Area								

Table 2: Displayed are the mean correlations with 95% credible intervals of the growth parameters for the three size variables. Parameters in bold denote

302 no overlap of the correlation coefficients with 0. Among-individual standard deviations are presented on the diagonal.

Juvenile predator cues show no strong effects on adult activity rates

Activity rates showed moderate repeatability from weeks 5-9 (*R* = 0.431 [0.327, 0.541]), demonstrating individual variation was stable. Activity generally decreased with time, though there were no clear effects of the treatments (Fig. 3). The control fish appeared to have lower activity rates, but differences among the treatment groups were inconsistent, with the combined treatment (i.e. Visual:Olfactory) being intermediate for females and the visual only intermediate for males. Also counter to our predictions, this effect was restricted to later trials – as such this effect needs to be interpreted with care.

The alternative character state model found variance increased in the last burst, while individual differences were largely maintained – consistent with the reported repeatability. This rank order consistency was very strong for females (for all combinations r > 0.71), though more modest for males (r = 0.37 - 0.57). There were additionally no among-individual correlations of activity rates with the growth parameters for body length in the alternative character-state model (correlation coefficients 0 ± 0.3 NS; see supplementary table).



Figure 3: Displayed are mean activity rates for treatment and temporal blocks with error bars representing the standard deviation of the credible distribution.

Discussion

We tested the effect of contradictory or incomplete information on informational state, and resulting developmental plasticity. Exposure to a predator altered adult size in all three of the measured variables, with the effect of the visual cues being stronger than the olfactory cues, indicating this was perceived as a more reliable source of information. However, the treatments did not have the predicted additive effect; we found no clear pattern of a larger response when the visual and olfactory stimuli were combined. The effect of predation on brain size was largely independent of overall body size, though appeared to lead to greater relative size in both sexes. In contrast, there was no effect of the predation treatments on activity rates, which also did not correlate with the growth parameters.

Effect of predation on growth

The visual cue had a strong effect on the development of guppies, increasing female brain size and reducing male body size. This indicates that the visual stimulus was perceived as a highly reliable source of information, which was not enhanced by the addition of olfactory cues. By contrast, the olfactory cue had a smaller effect, reducing male abdomen size, with no effect on females. This low reliability may help to explain the maladaptive effects when olfactory cues are presented alone (Ghalambor et al., 2015).

In manipulations of predator cues, there is an implicit assumption that the absence of cues (i.e. control) indicates that predators are at a low density and the environment is at low risk. Such information would likely be highly valuable to guppies from populations where larger predators are absent face other challenges owing to density regulations, competition and predation of juveniles by killifish (*Anablepsoides hartii*) (Travis et al., 2014). However, the absence of a cue may not be as reliable as the presence of a cue (Stamps and Bell, 2020). This is especially pertinent in cases such as predation, where events and exposure are infrequent and unpredictable, but clustered in time (Taborsky et al., 2020). Accordingly, we found that the visual stimulus had a strong effect – consistent with results over contextual time periods (Arteaga-Torres et al., 2020) – while the absence of complimentary olfactory cues did not reduce the response. Had we have provided a stimulus which indicated no-risk with greater certainty, a reduction of the response may have been more likely. For instance,

population demographics which reflect low predation environments can provide developmental cues of a low predation environment (Rodd et al., 1997).

Contrary to previous work showing that exposure to predation risk through development reduces somatic growth rates in guppies (Handelsman et al., 2013), none of our treatments affected the growth parameter (*k*). This meant that during the predator exposure phase of the experiment, there was no observed difference in size – divergence among treatments occurred after all individuals were returned to no-risk conditions. While compensatory changes in growth trajectories often occur in response to other environmental stressors (e.g. resources shortages or extreme temperatures) (Ali et al., 2003), predation in early ontogeny appeared to canalise later-life phenotype, while not affecting the measured phenotype during exposure. However, comparison to Handelsman et al. (2013) does require a caveat – while all effects of predation on growth rate were insignificant, best estimates were typically negative and we may not be able to reject a false-negative. Further, adult abdomen area appeared more sensitive than length, indicating effects of juvenile developmental stress may have reduced condition.

While studies of the evolved response of brain anatomy to predation pressure have provided variable results (Burns and Rodd, 2008; Kotrschal et al., 2017; Mitchell et al., 2020b; Reddon et al., 2018; Walsh et al., 2016), only one study has previously examined the developmental effects of predation risk (Reddon et al., 2018). These studies have focussed on differences in brain size after being standardised for body size, though results can reflect differences in starting size and growth of body size (Rogell et al., 2020). Here we explored growth in body and brain size as separate but correlated traits. Our results showed that while brain size was very highly correlated with body length and abdomen area at birth (r = 0.81 & 0.79 respectively), brain growth and adult size were less strongly correlated with the body size variables (r = 0.47 - 0.6), potentiating independent responses to predation risk.

Under exposure to predation, females had increased absolute brain size, while body size was largely unaffected – equating to an increase in relative brain size. This is likely an adaptive response, as previous experiments with brain size selected guppies have demonstrated benefits to antipredator behaviour and survival in large-brained females (Kotrschal et al., 2015; Kotrschal et al., 2013; van der Bijl et al., 2015). Males had smaller bodies when exposed to risk – consistent with evolved change from introduction

experiments (Reznick et al., 1990) – though brain size was unaffected. This also equates to an increase in *relative* brain size, and is consistent with results previously reported by Reddon et al. (2018). However, due to the lack of behavioural and survival advantages in the face of predation for males (Kotrschal et al., 2015; van der Bijl et al., 2015), and plasticity apparently limited to body size, differences in relative brain size among treatments seems unlikely to be adaptive for males.

No effect of juvenile predator cues on behaviour

We predicted a decrease in activity rates of predator exposed fish, so as to reduce vulnerability and encounter rates (Stamps, 2007), followed by convergence through time as fish updated their behaviour to match the standardised no-risk conditions. While predation risk affected adult size variables, and there was clear evidence for consistent amongindividual variation in activity, juvenile exposure to risk did not affect activity rates at 5 weeks old. Given this lack of an initial effect, we did not expect to see treatments affect temporal change through time. However, it was possible we would see divergence after the exposure to predation had ended. Pike cichlids target larger guppies (Johansson et al., 2004), while also reducing population sizes of killifish that prey on juvenile guppies (Travis et al., 2014). Accordingly, the perceived relative risk of our treatments may not have been high for five-weeks old fish. We observed some treatment level differences later in the experiment with control fish being more sedentary, though this did not seem intuitively meaningful as effects were counter to predictions and inconsistent among treatment groups. Further, contrary to the Bayesian models which predict convergence of phenotypes when animals are moved to standardised conditions (Stamps et al., 2018; Stamps and Krishnan, 2017), our character-state model revealed increased among-individual variance at nine weeks old.

Individual variation in activity was largely independent of the growth parameters. In addition to the lack of treatment differences, we found no correlations between activity rates and body length growth. Growth rates slow with age, and correlations may therefore be expressed at younger ages when animals are growing fastest – limiting hypothesised links of growth rates to earlier weeks. For our study, growth was fastest at 5-6 weeks old (burst 1) for males before slowing through time; for females, high growth rates were maintained through to the second burst of assays at 7-8 weeks old. Accordingly, we

complimented our analyses of treatment differences with a character-state model which allows us to address potential age-dependence of behaviour-growth correlations (Mitchell and Houslay, 2021). This analysis revealed no correlations between activity and growth parameters. Proposed correlations of life-history productivity and behaviour are built on an assumption that behaviours underly acquisition of resources (Biro and Stamps, 2008; Réale et al., 2010), though alternative allocation models may instead predict negative correlations due to energetic costs (Careau et al., 2008). These two factors are likely to be in balance (Laskowski et al., 2020), but as fish were fed routinely and were non-constrained by resources, the significance of activity to acquisition may have been dissociated.

Concluding remark

In this experiment, we aimed to test the effect of incomplete or contradictory information given by visual and olfactory cues of predation. Both cues affected the growth and morphology of guppies, reducing male body size and increasing female brain size, with effects of visual exposure appearing stronger. However, there was no evidence for a larger response when the stimuli were combined. Further, later life activity rates were also unaffected by exposure to predation. Together, our results indicate that visual and olfactory cues of predation were perceived as highly reliable sources of environmental information, while the absence of these cues as an indicator of low predator density was likely perceived as an unreliable source of information. Future work aiming to reduce the reliability of cues by altering frequency and/or concentration of stimuli, or working in systems where there is more evenly balanced perceived reliability of contrasting cues will be necessary to understand how informational state affects plasticity.

Data availability: Raw data and analysis code will be made available on publication

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Supplementary Table 1: Univariate activity model.

The following table corresponds to the activity analysis, which underlies Figure 3 in the main text.

				Est	SD[est]	Q2.5	Q97.5
	Control	Contol	BurstA	0.285	0.248	-0.206	0.761
	Visual	Contol	BurstA	0.450	0.268	-0.076	0.978
	Control	Olfactory	BurstA	0.241	0.303	-0.353	0.842
	Visual	Olfactory	BurstA	0.191	0.286	-0.377	0.749
s	Control	Contol	BurstB	-0.108	0.249	-0.593	0.383
ale	Visual	Contol	BurstB	0.418	0.264	-0.093	0.938
em	Control	Olfactory	BurstB	0.289	0.301	-0.301	0.878
	Visual	Olfactory	BurstB	-0.252	0.283	-0.805	0.300
	Control	Contol	BurstC	-0.554	0.255	-1.052	-0.050
	Visual	Contol	BurstC	0.118	0.265	-0.399	0.641
	Control	Olfactory	BurstC	0.029	0.303	-0.563	0.623
	Visual	Olfactory	BurstC	-0.468	0.284	-1.031	0.098
	Control	Contol	BurstA	-0.034	0.284	-0.590	0.517
	Visual	Contol	BurstA	0.163	0.250	-0.330	0.656
	Control	Olfactory	BurstA	0.400	0.215	-0.017	0.824
	Visual	Olfactory	BurstA	-0.129	0.259	-0.653	0.371
	Control	Contol	BurstB	-0.291	0.281	-0.856	0.259
les	Visual	Contol	BurstB	0.152	0.253	-0.344	0.644
Ba	Control	Olfactory	BurstB	0.070	0.217	-0.348	0.498
	Visual	Olfactory	BurstB	-0.022	0.255	-0.516	0.483
	Control	Contol	BurstC	-0.759	0.280	-1.317	-0.209
	Visual	Contol	BurstC	-0.445	0.253	-0.953	0.045
	Control	Olfactory	BurstC	-0.009	0.215	-0.430	0.414
	Visual	Olfactory	BurstC	-0.076	0.261	-0.594	0.435
		σ	ID	0.654	0.067	0.533	0.799
		σ	ID:Burst	0.378	0.045	0.291	0.468
		σ	е	0.645	0.020	0.607	0.686

Supplementary Table 2: Character-state model

The following is the table for the full output from the character-state model assessing covariances between the growth parameters for body length and activity rates at 5, 7 and 9 weeks old. Best estimates are given as the mean coefficient with the standard deviation and 95% credible intervals as quantiles.

				Est	SD[est]	Q2.5	Q97.5
	Female		S_0	2.034	0.012	2.010	2.057
	Male		S_0	1.985	0.012	1.961	2.010
	Female		k	-2.905	0.025	-2.954	-2.856
	Male		k	-2.677	0.022	-2.720	-2.633
	Female		S_{∞}	2.969	0.012	2.944	2.993
gth	Male		S_{∞}	2.792	0.008	2.776	2.808
leng		$ln(\sigma)$	int	-1.186	0.066	-1.314	-1.056
∕p		$ln(\sigma)$	Age	0.022	0.013	-0.002	0.046
Bo	le	σ	S0	0.069	0.009	0.052	0.089
	ema	σ	k	0.129	0.022	0.091	0.177
	Ĩ	σ	S_{∞}	0.065	0.010	0.047	0.087
	e	σ	S_0	0.075	0.010	0.059	0.097
	Mal	σ	k	0.123	0.020	0.087	0.166
	E	σ	S_{∞}	0.049	0.007	0.037	0.065
	Female		BurstA	0.300	0.128	0.049	0.550
	Male		BurstA	0.138	0.113	-0.087	0.361
	Female		BurstB	0.081	0.134	-0.179	0.343
	Male		BurstB	0.000	0.102	-0.203	0.197
>	Female		BurstC	-0.223	0.162	-0.537	0.096
ivit	Male		BurstC	-0.272	0.144	-0.556	0.006
Act	ale	σ	S_0	0.705	0.100	0.529	0.923
	emő	σ	k	0.734	0.101	0.558	0.951
	ŭ	σ	S_{∞}	0.901	0.117	0.698	1.155
	Ð	σ	S_0	0.656	0.096	0.486	0.865
	Mal	σ	k	0.579	0.090	0.418	0.772
		σ	S_{∞}	0.899	0.114	0.704	1.146
	Γ						
		S_0	k	-0.071	0.181	-0.412	0.287
		S_0	S_{∞}	0.279	0.167	-0.069	0.585
		k	S_{∞}	0.021	0.197	-0.358	0.408
		S_0	BurstA	0.008	0.171	-0.327	0.343
suo	S	k	BurstA	0.134	0.183	-0.234	0.473
lati	nale	S_{∞}	BurstA	0.006	0.181	-0.346	0.358
Jrre	Fen	S_0	BurstB	0.078	0.170	-0.258	0.408
ŭ		k	BurstB	0.068	0.180	-0.289	0.409
		S_{∞}	BurstB	-0.039	0.176	-0.374	0.307
		BurstA	BurstB	0.856	0.084	0.656	0.971
		S_0	BurstC	0.162	0.163	-0.165	0.469
		k	BurstC	0.274	0.166	-0.072	0.574

					1	
	S_{∞}	BurstC	-0.139	0.174	-0.469	0.200
	BurstA	BurstC	0.715	0.117	0.448	0.901
	BurstB	BurstC	0.803	0.094	0.583	0.942
	S_0	k	0.418	0.176	0.053	0.744
	S_0	S_{∞}	0.077	0.168	-0.255	0.392
	k	S_{∞}	-0.255	0.173	-0.565	0.110
	S_0	BurstA	-0.147	0.167	-0.466	0.183
	k	BurstA	-0.222	0.180	-0.557	0.146
	S_{∞}	BurstA	0.042	0.179	-0.309	0.388
S	S_0	BurstB	-0.176	0.168	-0.492	0.160
lale	k	BurstB	-0.102	0.190	-0.464	0.275
2	S_{∞}	BurstB	-0.129	0.177	-0.464	0.233
	BurstA	BurstB	0.579	0.153	0.246	0.833
	S_0	BurstC	-0.006	0.159	-0.315	0.299
	k	BurstC	0.016	0.176	-0.333	0.359
	S_{∞}	BurstC	-0.076	0.165	-0.392	0.251
	BurstA	BurstC	0.369	0.157	0.044	0.647
	BurstB	BurstC	0.570	0.143	0.262	0.816