



Amphibians breeding in refuge habitats have larvae with stronger antipredator responses



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Antipredator responses are a key determinant of the successful persistence of prey, and behavioural modifications are a frequent antipredator strategy. However, conspecific populations often inhabit heterogeneous environments. This can determine local adaptations, and might also induce variation in antipredator responses. Nevertheless, there is limited information on whether heterogeneity of predation risk among populations determines variation in antipredator response. Here we studied the fire salamander, *Salamandra salamandra*, a species that can breed in both surface streams and caves, habitats that are predator-rich and predator-free, respectively, and measured differences in antipredator responses across populations with different predation risk. We combined field surveys and laboratory experiments to understand the role of predation risk on the activity patterns of larvae, while measuring behavioural differences between populations. We reared larvae from different habitats in safe and risky conditions and tested their response to predator cues before and after rearing. In the field, predation risk was much higher in surface streams than in caves; larvae moved more in the absence of predators and when the light intensity was low. During laboratory experiments, larvae were less active if reared in risky conditions, but cave larvae showed a stronger response to risk than stream larvae. Therefore, larvae from sites without predators showed higher antipredator responses than those from risky habitats. This response fits the predictions of the risk allocation model, in which prey from habitats with a high background level of risk need to be active even when predators are present, to satisfy their energetic demands. Our findings show that antipredator behaviour may differ strongly between populations and stress the importance of integrating this variability in studies on predatory responses.

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Predators can affect prey through consumption, but also through nonconsumptive effects that alter the morphology, life history and behaviour of prey (Davenport & Chalcraft, 2013; Preisser & Bolnick, 2008; Winandy & Denoël, 2015), with major impacts on prey population dynamics (Davenport & Chalcraft, 2013). For instance, semiaquatic organisms with complex life cycles are able to assess the predation risk in a water body and modulate breeding activity by selecting habitats with fewer predators (Stav, Blaustein, & Margalit, 2000; Winandy, Darnet, & Denoël, 2015). Several studies have assessed the nonconsumptive effects of predation risk by evaluating how antipredator responses

enhance fitness and by measuring the costs and the trade-offs of such responses, which are keystones to understanding several ecological and conservation aspects (Amo, Lopez, & Martin, 2003; Blanchet, Bernatchez, & Dodson, 2007; Yorzinski et al., 2015). Such studies often use individuals of the same origin, without taking into account potential variation between populations (Blaustein, 1997; Hernandez & Peckarsky, 2014; Kishida, Trussell, Nishimura, & Ohgushi, 2009). However, local adaptations are common among populations that inhabit heterogeneous environments; thus spatial heterogeneity in predation pressure might induce variation in antipredator responses.

The reduction in activity levels under high predation risk is common antipredator behaviour (Kishida et al., 2009, 2011). Movements increase the risk of being detected by predators, but are often necessary to find resources such as food or partners, leading to a trade-off between reducing mortality and acquiring resources

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(Carlson & Langkilde, 2014). Some studies have underlined how individuals from populations living in risky habitats can display specific behavioural adaptations, which limit predation risk. In some cases, larvae of amphibians from habitats with high levels of predation risk (e.g. predatory fish) show a lower level of activity (e.g. moving less) and an even stronger reduction in activity rates in the presence of predators (Chivers, Wildy, Kiesecker, & Blaustein, 2001; Storfer & Sih, 1998). However, cases exist in which individuals from populations exposed to predators have higher activity levels, as this can improve their foraging and allow them to rapidly grow to a large size, thus reducing the period during which individuals suffer predation risk (Urban, 2007). In practice, anti-predator adaptations can be highly variable, suggesting that knowledge of how the heterogeneity of predation risk among populations and habitats determines the variation in antipredator response between populations is incomplete.

Predation risk is variable over space and time, and this variability is crucial in determining how prey respond to risk (Ferrari, 2014). As a consequence, the background level of risk is an additional factor potentially affecting the outcome of prey responses, as it can influence the readiness of a species to respond to predators (Ferrari, 2014; Ferrari, Crane, Brown, & Chivers, 2015). For instance, prey inhabiting habitats with high level of risk can decrease their vigilance and be particularly active during brief periods of safety (paradox of risk allocation hypothesis: Ferrari, Sih, & Chivers, 2009; Lima & Bednekoff, 1999). Furthermore, the background level of risk can strongly modify the overall activity level of prey, and this influences the outcome of their interactions with predators, with increase or decrease of survival depending on the predator type (Ferrari et al., 2015).

The fire salamander, *Salamandra salamandra*, is a usually epigeous taxon that breeds in streams, but some populations have also colonized subterranean environments in which larvae successfully complete their development (Manenti, Denoël, & Ficetola, 2013). On the one hand, subterranean environments can provide advantages, such as more stable environmental conditions and limited predation risk. On the other, salamanders in these habitats face major challenges, such as food scarcity (Manenti, Pennati, & Ficetola, 2015). Evidence suggests that local adaptations to underground environments are present, with cave populations also showing a higher behavioural plasticity than surface stream populations. This plasticity allows them to modulate activity levels in relation to environmental conditions and availability of prey, thus helping the colonization of such challenging habitat (Manenti, Denoël, et al., 2013; Manenti & Ficetola, 2013).

In this study, we combined field surveys and laboratory experiments to assess whether antipredator responses of salamander larvae differ between populations facing different levels of predation risk. First, we evaluated in the field whether movement is reduced in habitats with a higher predation risk. Second, we reared larvae from populations with a different level of predation risk (caves: absence of aquatic predators for salamander larvae; streams: presence of predators), and tested (1) whether larvae reared under risky conditions or in the presence of predatory cues are less active and (2) whether the response to risky conditions is similar or differs between populations that experience different predation risks.

METHODS

Field Surveys

We studied fire salamander populations from Lombardy (northwest Italy; approximately 45°48'N, 9°02'E). In this area, the fire salamander is ovoviviparous and usually produces larvae in

streams, but also often in caves (Supplementary Fig. S1). All these salamander larvae are fully aquatic and have external pairs of gills. Adult females actively enter caves and select them as breeding sites; no larvae in the chosen sites could have drifted into this environment from superficial waters, because all sites are springs that receive water exclusively from the subterranean aquifer (Manenti & Ficetola, 2013; Manenti, Ficetola, Bianchi, & De Bernardi, 2009). In these underground habitats, salamander larvae successfully grow and metamorphose (Manenti, Ficetola, Marieni, & De Bernardi, 2011).

We surveyed 52 sites (15 cave pools and 37 outdoor springs and stream pools; Fig. S1) between February 2014 and June 2015 during the day and at night. We visited each site twice and the same observer performed all the surveys. On all the sampling occasions, we recorded the number of active salamander larvae, i.e. the number of larvae visible from the pool border in 5 min visual surveys. To do this, we approached the pools in the dark and once at the border of the pool, we shone a torch on it (Petzl Ultra Vario) and directly counted the larvae. Subsequently, we estimated the total number of larvae using two successive removal samplings with a fine mesh net and applying the removal method (Chao & Chang, 1999). Larvae were released at their place of capture immediately after the census. We then calculated the frequency of active larvae as a proportion of the total number of larvae. As environmental variables, we recorded the maximum light intensity (illuminance) on the pool surface using a CEM DT8820 lux meter (CEM-instruments, Shenzhen, China), and estimated the biomass of predators (dragonfly larvae; g/m²) through pipe sampling (diameter: 25 cm; Dodd, 2010; see Limongi, Ficetola, Romeo, and Manenti (2015) for additional details on invertebrate samplings). All the studied pools were devoid of fish and we excluded from the analyses a few surface sites ($N = 5$) in which we found native crayfish, *Austropotamobius italicus*.

Laboratory Experiments

We performed behavioural experiments to assess the variation in movement between salamander larvae born under different risk conditions in underground and surface streams. We collected larvae at developmental stage 1 (newborns: well-developed tail fin and the tip of the fin bluntly rounded; Juszczuk & Zakrzewski, 1981) from underground pools (67 individuals from six sites) and neighbouring surface pools or slow-running water streams (96 individuals from eight sites). Larvae were individually maintained at a mean temperature of 18 °C, exposed at the natural photoperiod, in 10 × 11 cm perforated (diameter of perforations: 2 mm) transparent plastic containers placed in six independent water-filled blocks (i.e. plastic containers of 40 × 50 cm; water depth: 5 cm). Larvae were randomly assigned to three rearing treatments, with two blocks per treatment (each containing 12 larvae). The rearing treatments were a control (absence of dragonfly larvae), predator without contact ('no contact') and predator with contact ('contact'). Under the predator without contact treatment, two dragonfly larvae, *Cordulegaster bidentata*, were free to move within the block. Salamanders could receive dragonfly cues through the perforated walls of their container, but were protected from predation or direct contact. In the predator with contact condition, a dragonfly larva was placed in the container of the larva (see Ethical Note below), twice weekly, for 30 s. All *C. bidentata* larvae used were at the premetamorphosis stages (mean length ± SE = 31.8 ± 0.4 mm) and are major predators of salamander larvae within the study area (Manenti, Siesa & Ficetola, 2013). *Salamandra salamandra* larvae were fed ad libitum every 2 days with *Chironomus* sp. larvae. Chironomids were also provided ad libitum to *C. bidentata* larvae for a total of 8–10 prey per week. Salamander larvae were kept without food for 3 days before

we performed behavioural tests to get the same level of satiety. This absence of feeding does not affect salamander success and occurs in the wild (Limongi et al., 2015). The body size (total length) did not differ between salamander larvae in the caves or streams, or between rearing conditions (at the beginning and at the end of the experiment, *t* tests showed for all $P > 0.2$).

During behavioural tests, we assessed the effect of the presence of predator cues on salamander movements. These tests allowed us also to determine the effect of the site of origin (surface streams versus cave) and rearing conditions (predator and predator-free environments). Predator cues were obtained by maintaining six *C. bidentata* larvae for 24 h in 1.5 litres of dechlorinated tap water; the water was aliquoted (1 ml) and immediately stored at -20°C until use for behavioural tests, following the procedure of Epp and Gabor (2008). As control cues, we used 1 ml of tap water. Behavioural tests started 3 days after salamander collection, and were repeated after 45 days of rearing; the identity of the larvae to be tested in each trial and the treatment order were randomly selected, until each larva was tested twice with the predator and twice with control cues. Both tests were repeated 3 days after salamander collection and at 45 days of rearing. During the behavioural tests, each larva was individually placed in a 13.5×18.3 cm plastic container filled with 5 cm of water under daylight conditions with an average lux intensity of 500 lx and was allowed to acclimatize for 3 min. At the beginning of the test, 1 ml of water with the test cues was cautiously placed with a pipette on the opposite side of the arena by the same person. The trials lasted for 7 min. The entire observation tank was video-recorded (Panasonic SDR-S7, Bracknell, U.K.) from above during all the trials.

We then used the Noldus Ethovision XT10 video-tracking software (Noldus Information Technology, Wageningen, The Netherlands) to measure the distance moved by the individuals during each trial. This software automatically detected salamander larvae on the basis of their contrast with the background and gave them a position over time (25 images/s over 7 min/trial in our setting; Delcourt, Denoël, Yliff, & Poncin, 2013). By using this automated method, it was possible to obtain accurate locations of salamander larvae across time without any observer effect (Delcourt et al., 2013). As a behavioural parameter, we considered the total distance moved by the larva, since it represents a valuable indicator of the activity rate of amphibian larvae (Denoël et al., 2010; Uiblein, Durand, Juberthie, & Parzefall, 1992; Uiblein et al., 1995) and strongly influences feeding performance, as larvae that move greater distances have better feeding success (Manenti, Denoël, et al., 2013).

Statistical Analyses

We used a linear mixed model (LMM) to assess whether the activity of larvae in the field was related to predation risk. The frequency of active larvae (arcsine square root transformed) was the dependent variable. As an independent variable, we used the log-transformed biomass of predators. We included the log-transformed incidence of light (activity levels are strongly affected by light conditions; Manenti, Denoël, et al., 2013) and habitat typology (cave/stream) as additional covariates. Site identity was included as a random factor. We used marginal and conditional R^2 (R^2_{m} and R^2_{c} , respectively) to assess the fit of the LMM. The R^2_{m} represents the pure effect of fixed factors, whereas R^2_{c} is the variance explained by the entire model (Nakagawa & Schielzeth, 2013).

We used LMM to assess the factors that determined the behavioural responses of larvae in laboratory experiments to test and rearing conditions. The total distance moved (log-transformed)

was the dependent variable. We considered four fixed independent factors: the origin of the larvae (cave/stream); the rearing conditions (control, no contact, contact), the presence of predator cues during the behavioural tests and the time of the test (the beginning of the experiment versus after 45 days). The origin was intended to test for differences between possible ecotypes; rearing conditions were intended to test for the influence of growing in different types of risky habitats and the test conditions were assumed to test for the immediate response to risk cues. We also tested two-way interactions. The final model included all the fixed factors and all the significant interactions. We used orthogonal contrasts to evaluate the significance of differences between the three rearing conditions (controls versus with predators; no contact versus contact). All our models included larva identity, site of origin and rearing block as random factors. Among random factors, rearing block explained a very small amount of variation, as removing block from the model slightly decreased Akaike information criterion (AIC) values (complete model: 1645.8; model without block: AIC = 1644.7). We present the results taking into account rearing block, but results remain identical if this factor was not included in analyses.

The sample size was not homogeneous among populations; therefore, in mixed models, the degrees of freedom were approximated (Satterthwaite, 1946). Analyses were performed using the lme4 and lmerTest packages in R (Bates, Maechler, Bolker, & Walker, 2014; Kuznetsova, Brockhoff, & Christensen, 2015).

Ethical Note

The study was approved by the ethical committee of the Lombardy Region Authority and was authorized as complying with the regional law 10/2008, p.n.: F1.2013.0002091. The survival rate of larvae was 70.4%, being much higher than that observed in natural populations (Limongi et al., 2015). All the larvae were released at their site of origin at the end of the study, following the recommendations of the permit. All individuals were checked daily and fed every 2 days. The aim of this study was to evaluate only the nonconsumptive effects of dragonfly larvae on salamander larvae by observing their behaviour. Consequently, care was taken to plan the experiment accordingly and, thus, to avoid any larvae being harmed. To this end, dragonfly larvae were placed for only 30 s in the tank of salamanders, thus preventing any predation. Moreover, we stopped them from extending their mouthparts and were always ready with tweezers to block snaps. This design was needed because one of the aims of the study was to compare indirect and direct contact with dragonfly larvae (see also Winandy & Denoël, 2013b, 2015).

RESULTS

The mean abundance of predators in streams was 1.06 g/m^2 ($\text{SE} = 0.48$; mainly dragonflies of the genus *Cordulegaster*), whereas we never captured dragonfly larvae in caves. Overall, a model including light, abundance of predators and type of site explained the variation in the activity level in natural conditions very well ($R^2_{\text{m}} = 0.60$; $R^2_{\text{c}} = 0.67$). The proportion of active larvae decreased at high light levels ($F_{1,50.9} = 113.6$, $P < 0.001$) and in sites with more predators ($F_{1,51.5} = 7.6$, $P = 0.008$). When taking into account the effect of light and predator abundance, the frequency of active larvae did not differ between caves and streams ($F_{1,55.6} = 0.1$, $P = 0.753$; Fig. 1).

In laboratory trials, the total distance moved ranged between 0 and 431.93 cm (mean = 47.81, $\text{SD} = 61.15$). The total distance moved decreased after 45 days of rearing (Table 1). The decrease was stronger in larvae reared under risky conditions than in controls (orthogonal contrast: $F_{1,826.8} = 7.45$, $P = 0.006$; Fig. 2a),

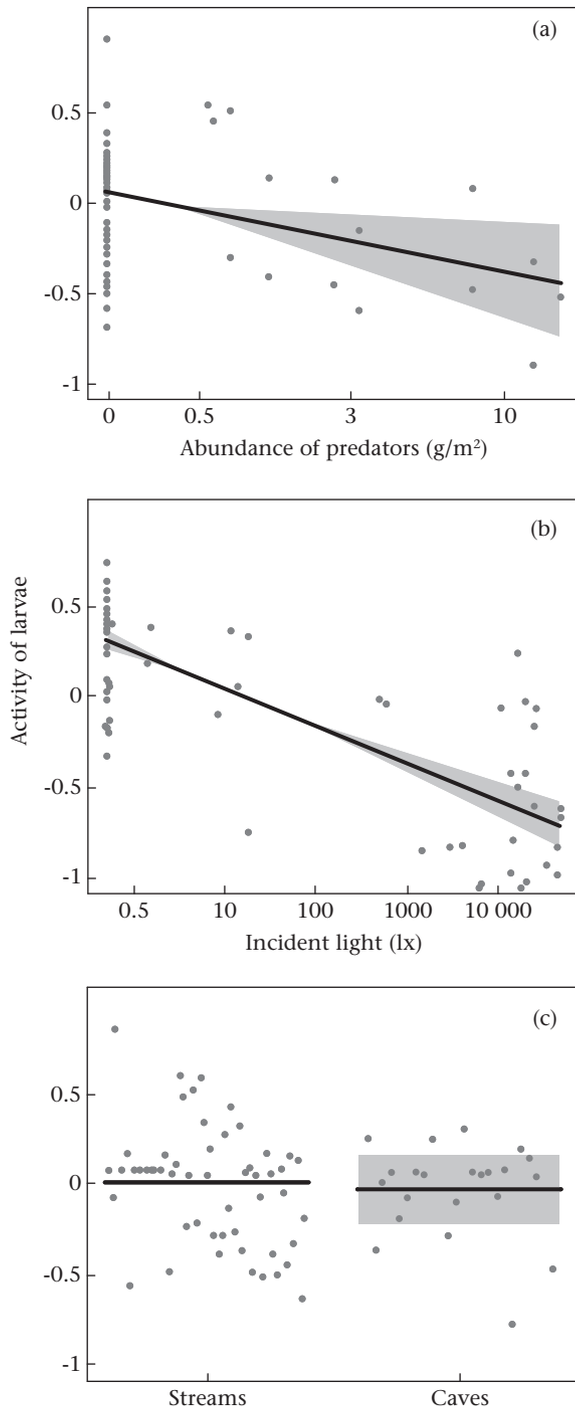


Figure 1. Contrast partial regression plots, showing the relationships between the proportion of active fire salamander larvae under natural conditions (arcsine square root transformed) and environmental variables: (a) illuminance, (b) predator abundance and (c) site. Shaded areas represent the 95% confidence bands. Plots were built using visreg 2.2 (Breheny & Burchett, 2015).

whereas no difference was detected between rearing in contact with the predator and rearing with the predator but without contact ($F_{1,847.6} = 2.34$, $P = 0.126$). Only larvae from caves showed a reduced activity in the presence of predator cues ($F_{1,706.7} = 9.66$, $P < 0.01$; Fig. 2b). Furthermore, the activity of larvae from caves decreased with time more than that of larvae from streams ($F_{1,837.3} = 4.33$, $P = 0.03$; Fig. 2c).

Table 1
Variables influencing the activity of fire salamander larvae

Variable	<i>F</i>	<i>df</i>	<i>P</i>
Origin (cave/stream)	0.17	1,14.8	0.690
Time of test	16.36	1,831.0	<0.001
Rearing conditions	0.05	2,2.1	0.950
Test conditions	0.43	1,708.2	0.511
Origin * test conditions	9.66	1,706.7	0.002
Origin * time of test	4.33	1,837.3	0.038
Rearing conditions * time of test	4.99	2,836.8	0.007

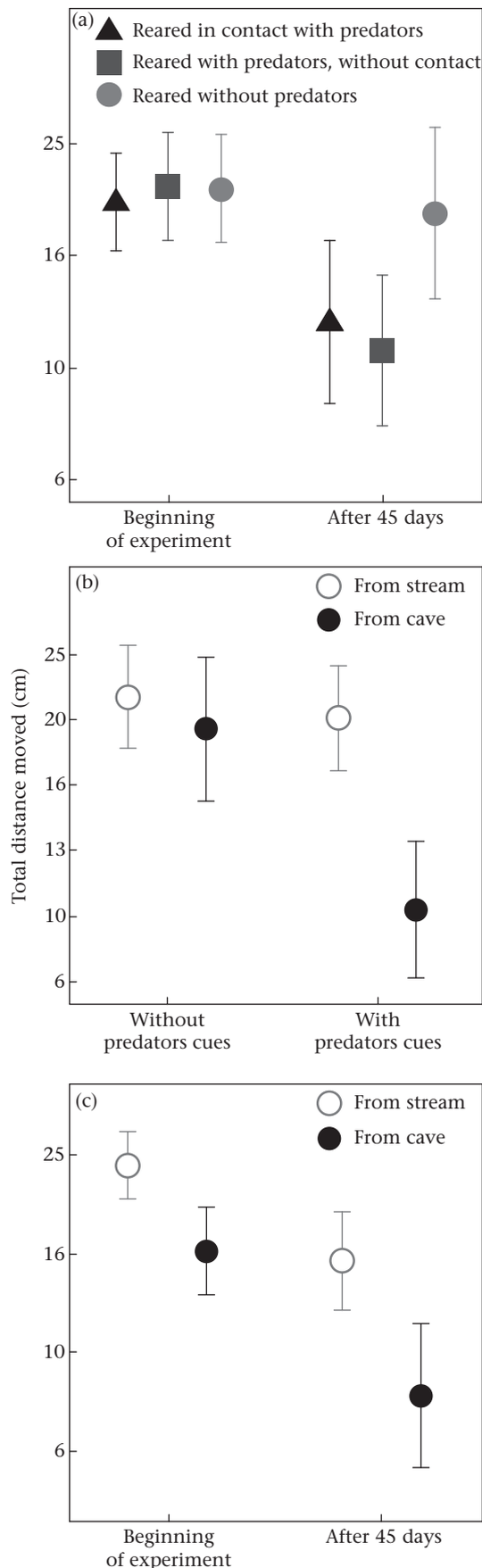
Variables influencing the activity (i.e. total distance moved) of fire salamander larvae: results of mixed models assessing the effect of origin, rearing conditions and test conditions (with or without predator cues) on the movement of larvae. Only significant interactions are included in the model. The degrees of freedom are approximated following Satterthwaite (1946); thus, some are not integers.

DISCUSSION

The prediction of species interactions across heterogeneous habitats is a major challenge for zoologists and ecologists. Within a single species, adaptations can change across neighbouring localities (Caspers, Steinfartz, & Krause, 2015; Krause & Caspers, 2015), given that conspecific populations that exploit different habitats are subject to different selective pressures (Godsall, Coulson, & Malo, 2014). Longstanding ethological theory predicts that prey should show strong responses to threats, such as decreasing overall activity or escaping to reduce predation risk (Barnett & Richardson, 2002; Carlson & Langkilde, 2014; Lima, 1998). However, apparently opposite strategies, such as risky behaviours, might evolve in certain predator-rich habitats (Urban, 2007), and the risk allocation model has been proposed and developed to explain such contrasting strategies (Ferrari et al., 2015; Lima & Bednekoff, 1999). The model predicts that, if predators are not always present, prey should show strong antipredator responses, while they are expected to increase their foraging effort and activity levels during low-risk situations. Conversely, living in risky conditions, in which predators are always present, can determine a limited antipredator surveillance, because prey may need to forage even when predators are present (Ferrari et al., 2009). Our study is in agreement with this model, as salamander larvae from populations breeding in the safer habitats (i.e. caves) showed a stronger antipredator behaviour (i.e. activity reduction) than those from populations in predator-rich habitats (i.e. streams).

The combination of field and laboratory observations allowed us to obtain a complete picture of how antipredator behaviour is modulated as a function of predation risk. In the field, light conditions and predation pressure both affected the movement patterns of fire salamander larvae: movements were more limited in the presence of predators and of light, i.e. when there is more risk of being detected by predators. In caves, darkness is constant and predators are absent. These environments thus provide salamander larvae with a habitat with the lowest predation risk (Uiblein et al., 1992).

Larvae reared with dragonflies moved less than control larvae. Recent studies have proposed that direct contact with a potential predator can be perceived as a threat by aquatic newts more than the simple presence of cues (Winandy & Denoël, 2013b). In direct contact, salamander larvae might also experience a closer disturbance or perception of potential risk from the dragonfly, increasing their caution and decreasing their movement. The treatment involving no direct contact with the predator was thus performed to test for the occurrence of habituation or different levels of threat perception, as have been observed in other study systems (Winandy & Denoël, 2013a). Nevertheless, we did not detect any difference between conditions involving or not involving direct



contact, indicating that dragonfly larvae were perceived as a risk, even when they were not in direct contact with the salamanders in our study system. This result may be related to the fact that the duration of exposure was not the same: in the indirect condition, salamanders were continuously exposed to the risk of predation, while in the direct condition, the risk of predation was very short in order to prevent the consumption of test larvae. Further tests should therefore focus on possible habituation effects. However, it is interesting that two very short direct exposures to predators per week led to the same effect as permanent indirect exposure, underlying the strong impact of direct contact.

The behavioural responses of cave and stream larvae differed considerably, as cave larvae showed the strongest reduction in activity in the presence of predators during rearing, and in the presence of predator cues during tests. Stream larvae did not move less in the presence of predator cues, indicating that individuals from the riskier environment assume the riskier behaviour (Urban, 2007). These results may be explained by Ferrari et al.'s (2009) risk allocation hypothesis: in low-risk habitats, when predators are absent or rare, prey can feed during relatively long safe periods and thus respond strongly to short and occasional presence of predators. In contrast, if conditions are highly risky with abundant predators, prey need to be active and forage even though predator cues occur. Streams are riskier for other parameters, such as desiccation before metamorphosis, whereas underground sources in our study area are always wet (Manenti, 2014). In ephemeral streams, with the constant presence of predators, a cautious behavioural strategy might cause an excessively long larval phase, also involving a higher risk of stream desiccation before metamorphosis. Conversely, the cost of reduced activity might be lower in cave pools, where predators occur only occasionally and delayed metamorphosis does not necessarily increase mortality. The relative fitness of individuals with risky and prudent antipredator strategies might thus be context dependent and be strongly influenced by the interaction between predator presence, predator type and the abiotic features of the breeding sites. Moreover, populations may also diverge in growth rate, thereby influencing the effectiveness of different antipredator strategies (Laurila, Pakkasmaa, & Merilä, 2006).

The predation risk differed considerably between surface and underground sites: caves were devoid of potential predators, whereas streams were often inhabited by dragonfly larvae. Dragonflies are efficient predators of amphibian larvae and can have a strong impact on the density of salamanders (Manenti, Siesa, et al., 2013), with a high impact on population dynamics (Drake, Anderson, Smith, Lohraff, & Semlitsch, 2014; Kishida et al., 2009). Conversely, caves lack interspecific predators, but the limited food resources of underground habitats delay larval development (Limongi et al., 2015). When selecting breeding sites, females thus face two potential strategies: a risky strategy in streams and a more prudent strategy in caves.

In the populations studied here, it is not known whether differences between cave and stream larvae are mostly determined by local adaptations, or whether nongenetic maternal effects play a role. Responses to predators generally depend on two mechanisms: exposure to different background levels of risk (Ferrari, 2014) and associative learning that allows prey to associate novel stimuli with danger (Mathis, Ferrari, Windel, Messier, & Chivers, 2008).

Figure 2. Effect of experimental conditions and origin on the activity (i.e. distance moved) of fire salamander larvae: analysis of experimental data. (a) Effect of rearing conditions and time of the test; (b) effect of salamander origin and test conditions; (c) effect of time of the test and salamander origin. All the interactions shown in the plots are significant (see Table 1). Error bars are two standard errors of the mean.

Selection is predicted to favour early learning in development, and it has been shown that even embryonic conditioning affects larval antipredator responses in amphibians (Ferrari et al., 2015; Mathis et al., 2008). Fire salamanders are ovoviviparous; thus embryonic exposure is impossible. Nevertheless, although we caught the larvae shortly after birth, we cannot exclude an influence of the very first posthatching experience to which larvae were exposed in their original habitat. In any case, the strong response to the predator in individuals from cave populations suggests some contact between these predators and the fire salamanders breeding in caves. Because local adaptations often reduce the ability to recognize or respond to predators that the populations have not encountered during recent evolutionary history, it is likely that gene flow occurs between salamanders that breed in caves and in streams. Indeed, individuals that metamorphose in caves usually leave them and can then potentially breed with individuals originating from the surrounding streams or springs, even though assortative mating might limit the gene flow between stream- and cave-breeding females, thus allowing the persistence of local adaptations (Caspers, Junge, Weitere, & Steinfartz, 2009). Further population genetics studies are necessary to determine the potential differences between cave and stream populations.

The heterogeneity of predation pressure across environments can determine strong intraspecific variation in behaviour. Such variability can be a key factor favouring the colonization of novel habitats, and can also arise between nearby populations living in contrasting habitats. However, studies on antipredator responses often focus on a single population. As antipredator responses may show contrasting patterns among habitats, we stress the importance of including between-population variation in this kind of study.

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Supplementary Material

Supplementary material related to this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.anbehav.2016.06.006>.

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