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Changes in predator exposure, but not diet induce phenotypic plasticity in scorpion venom

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Figure 1. Averaged relative venom profiles for each of the four treatments taken: prior to treatments (A, t = 0 days), after the treatments ended (B, t = 49 days), and again after a recovery period (C, t = 70 days) divided into 14 fractions. Relative absorbance measures the absorbance units at 280 nm of any point along the venom profile relative to the point of maximum absorbance in the profile. Venom profiles obtained from scorpions subjected to the pressure for defensive venom (mouse exposure) treatment are given in dark blue (+ pressure for offensive venom, live cricket prey) and light blue (- pressure for offensive venom, dead cricket prey); profiles obtained from scorpions not subjected to this treatment are given in red (+ pressure for offensive venom) and orange (- pressure for offensive venom).



Figure 2. Loadings for principal components 1 and 2, separated into loadings for each scorpion (A) and loadings for each venom fraction (B, 14 total peaks) with fraction number indicated at the tips of each arrow. For clarity and consistency, colours are as per the curves presented in Figure 1; pressure for defensive venom treatment is indicated in dark (+ pressure for offensive venom) and light (- offensive pressure for offensive venom) blue, while data from scorpions not subjected to the pressure for defensive venom treatment are in red (+ pressure for offensive venom) and orange (- pressure for offensive venom). PC1 and PC2 described 53.6% and 25.4% of the overall variation, respectively.

238x132mm (96 x 96 DPI)



1	Changes in predator exposure, but not diet induce phenotypic plasticity in scorpion venom
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8	
9	Abstract
10	
11	Animals embedded between trophic levels must simultaneously balance pressures to deter
12	predators and acquire resources. Venomous animals may use venom toxins to mediate both
13	pressures, and thus changes in this balance may alter the composition of venoms. Basic
14	theory suggests that greater exposure to a predator should induce a larger proportion of
15	defensive venom components relative to offensive venom components, while increases in
16	arms races with prey will elicit the reverse. Alternatively, reducing the need for venom
17	expenditure for food acquisition, for example due to an increase in scavenging, may reduce
18	the production of offensive venom components. Here, we investigated changes in scorpion
19	venom composition using a mesocosm experiment where we manipulated scorpions'
20	exposure to a surrogate vertebrate predator and live and dead prey. After six weeks, scorpions
21	exposed to surrogate predators exhibited significantly different venom chemistry compared to
22	naïve scorpions. This change included a relative increase in some compounds toxic to
23	vertebrate cells, and a relative decrease in some compounds effective against their
24	invertebrate prey. Our findings provide, to our knowledge, the first evidence for adaptive

- plasticity in venom composition. These changes in venom composition may increase thestability of food webs involving venomous animals.
- 27

28 1 Introduction

29 Interspecific arms races are ubiquitous in ecological communities, and generally involve 30 reciprocal selection pressures that drive the evolution of adaptations and responses between 31 interacting organisms. Organisms investing in traits mediating these arms races need to balance the fitness benefits of winning the race against the cost of maintaining those traits [1]. 32 33 Some organisms defend themselves in arms races using chemical toxins, and these toxins are 34 often produced in low quantities unless induced by exposure to natural enemies to minimize 35 the cost of maintaining unnecessary defences [2]. Similarly, when predators are exposed to 36 prey with varying defensive adaptations, they may develop inducible chemical weapons [3, 37 4]. In venomous animals, the same delivery apparatus evolved for prey capture – such as fangs or a stinger - can also be used to inject chemicals to deter enemies [5], and this dual-38 39 purpose nature of the delivery apparatus also extends to the chemistry of the venom itself. 40 Animals generally need to balance arms races involving both predators and prey, and these 41 arms races drive the evolution of venom chemistry in both offensive and defensive contexts 42 [5]. However, physiological differences between predators and prey may necessitate different 43 toxins, and specificity of venom toxins to particular groups of animals has been identified in 44 many venom-users [6-11]. For example, sodium channel blocking α -toxins in scorpions contain three separate subtypes of toxins that are effective against the voltage-gated sodium 45 channels of mammals only, insects only, and both [6]. The whole venom mixture can be 46 47 thought of as a cocktail of these different toxins, but whether the 'recipe' for this cocktail is 48 fixed or can exhibit plasticity in response to different environments and predator/prey 49 interactions remains unclear [12].

50	Broadly speaking, plasticity will be favourable when it enables an organism to have
51	higher fitness across multiple environments, or within a variable environment [13].
52	Theoretical work has shown that plasticity can be selected for when: (a) populations are
53	exposed to multiple environments or variability within an environment, (b) environments
54	produce reliable cues, (c) different phenotypes are favoured in each environment, and (d) no
55	single phenotype exhibits superior fitness across all environments [14-16]. Both plants and
56	animals can in principle exhibit 'induced' plasticity to calibrate their defences in response to
57	species interactions [17]. In this context, plasticity may provide a way of saving costs
58	associated with defences, allowing resources to be allocated towards growth and reproduction
59	instead [18]. For example, theory suggests that costly plant defensive compounds should be
60	constitutive (i.e. permanent) where the probability of herbivory is high, while an induced
61	defence in response to attack is optimal when the probability of herbivory is low but the
62	threat of injury from an attack is high [19, 20] Though less well-understood, induced
63	offensive traits enable a predator to capture certain prey more efficiently via plastic change in
64	response to cues signalling the presence of that prey [21]. Induced offences are more
65	favourable when a consumer can benefit from adapting to multiple resource (prey) species
66	with a variety of defences, or resources that can exhibit variable levels of a defence [22, 23].
67	For example, plastic induced 'offensive traits' can be seen in feeding-morphologies, such as
68	in snails from the genus Lacuna, which change the shape of their teeth to suit their prey [24],
69	and Nephila pilipes spiders can plastically modify the composition of their silk chemistry in
70	order to vary the architecture and physical properties of their webs to catch different prey
71	[25].
72	In venom-users, the high cost of chemical warfare has selected for a range of

behavioural 'venom-metering' strategies, and these plastic behaviours are used to minimize
 the quantity of venom delivered. Spiders, for instance, may evaluate venom resistance in prey

75 based on olfactory cues and use their venom accordingly [26]. A choice of whether or not to 76 envenomate at all has been shown, based on the relative size and threat posed by the target 77 [27, 28]. Once the decision to envenomate is made, a range of other cues can influence the 78 delivery and volume of venom to minimize venom-use across venomous taxa [12]. By 79 employing similar cues, a venom-user is able to modify the 'recipe' of its venom cocktail, 80 thereby optimizing the fitness benefits of its costly venom in different environments 81 exhibiting differences in densities and types of predators and prey [4]. Analogous to induced 82 non-injected defences, a fixed, constitutive venom 'recipe' may be more favourable in 83 environments with higher rates of predator attack and lower variability in predator type, while 84 a plastic 'recipe' may be more favourable where rates of predator or prey encounters, or 85 predator or prey types, are variable. However, to our knowledge neither induced defensive 86 toxin production nor induced offensive toxin production have been demonstrated in 87 venomous animals.

Here, we present an experimental exploration of induced plasticity in the composition 88 89 of venom produced by scorpions in response to perceived predation risk, and reduced need 90 for venom-use for prey capture. We test the hypotheses that, if induced plasticity of venom 91 composition is exhibited by a model venom-user, higher predation risk will lead to higher 92 relative production of predator-active toxins; and that relative production of prey-active 93 toxins will increase in response to a prey-type that requires greater venom-expenditure to 94 ensure a meal. We did this by manipulating exposure to a surrogate predator and access to 95 live or dead prey, and evaluating changes in the relative concentrations of prey-specific 96 toxins, predator-specific toxins, and general venom compounds. To investigate whether 97 manipulated rates of predator-prey interactions would elicit the plastic changes in venom 98 composition we used the Australian rainforest scorpion *Liocheles waigiensis* (Gervais) (Scorpionoidea: Hemisorpiidae). Our next goal was to evaluate the effects of the predator-99

100	specific toxins present in L. waigiensis venom on mammalian cells and the prey-specific
101	toxins on crickets that represent their invertebrate prey. Finally, we evaluated whether the
102	relative proportions of vertebrate-toxic venom components would increase in response to a
103	higher rate of vertebrate predator-interactions, and whether the relative proportions of
104	invertebrate-toxic components would decrease in response to a lower need for venom in food
105	consumption, through increased scavenging behaviour.
106	
107	2 Methods
108	(a) Model organism
109	The Australian rainforest scorpion Liocheles waigiensis (Gervais) (Scorpionoidea:
110	Hemisorpiidae) used in our experiments is a common species found in the wet tropics of Far
111	North Queensland. L. waigiensis is a generalist predator of invertebrates, including crickets,
112	and is in turn preyed upon by a range of invertebrate and vertebrate predators [29]. Scorpions
113	sourced from rainforest areas around Cairns were individually held in $170 \times 110 \times 50 \text{ mm}$
114	650 ml plastic containers with one stone and moist organic soil (300 ml Searles Premium
115	Potting Mix brand potting mix) to provide a suitable microclimate for the animal and to aid
116	with moulting. These containers were randomly sorted and stacked two high, in two
117	Wisecube WGC-450 temperature and humidity chambers at 28 °C on a 14/10 light/dark
118	cycle. Relative humidity was maintained at 70%, and after 3 weeks of treatments all
119	scorpions were moved to new containers containing freshly autoclaved soil to reduce fungal
120	growth. All scorpions were maintained in the controlled environment for no more than 5 days
121	prior to the first venom extraction.
122	

123 (b) Experimental treatments

124 Given the general predation of invertebrates by L. waigiensis in the wild, we used the 125 common house cricket Acheta domesticus L. (Insecta: Orthoptera) as a surrogate prey 126 species. To simulate a vertebrate predator sometimes encountered by L. waigiensis in the 127 wild, a frozen feeder mouse, *Mus musculus* L. (Rodentia: Muridae), was taxidermied by 128 skinning, stuffing with cotton wool, and articulation with wire, then used as a simulated 129 model vertebrate predator (as in Digweed and Rendall [30]). We used a 2 x 2 factorial design 130 in which without and with pressure for induced offensive venom production (presence of live 131 versus dead prey) was crossed with pressure for defensive venom production (simulated 132 predator exposure). The number of replicates, accounting for scorpions that died during the 133 treatments and were therefore excluded from the analysis, were as follows: 15 (pressure for 134 offensive + defensive venoms), 14 (pressure for offense venom), 14 (pressure for defensive 135 venom), 13 (control).

136 For the prey treatment, scorpions were each fed either a live (pressure for offensive venom) or dead cricket once per week. Live crickets were purchased 1-2 days prior to each 137 138 feeding, and were killed by freezing for approximately 12 hours. In doing so, the quality of 139 the diet was identical for both groups, but the pressure to use venom to obtain a meal was not. 140 Our taxidermied mouse was used to provoke defensive stings from scorpions in the defensive 141 venom pressure treatment three times a week, except for the first week to allow for 142 acclimation. The mouse was used to continuously probe on the cephalothorax of defensive 143 pressure treatment scorpions for 30 seconds. This stimulus readily stimulated anti-predator 144 responses in the scorpions, including alert and threat postures (with chelae extended and 145 open, and metasoma erect), grappling, pinching, stinging, squirming, and retreat [31, 32]. To 146 ensure that scorpions excluded from the defensive pressure treatment were otherwise equally 147 handled and exposed to laboratory conditions, the containers of these scorpions were opened 148 and exposed to laboratory conditions for 30 seconds. Six weeks after commencement of the

149	experiment, scorpions were subjected to the control (no offensive, no defensive pressures)
150	treatment for one week before venom was extracted. Although this may have diminished the
151	measured effects of the with-pressure treatments due to relaxing of any induced response, a
152	brief recovery time was necessary to ensure sufficient volume of venom had recovered to
153	perform chemical analyses.
154	
155	(c) Venom extraction
156	We ran the treatments for 42 days so that the experiments lasted twice as long as the venom
157	regeneration time of 21 days, according to previous analyses [29]. Venom was first extracted
158	within 5 days of collection, and then again after a week of rest, following the end of the
159	experiment, 49 days later. Venom was then extracted from all scorpions a third time, 21 days
160	after the experimental treatments ceased, to assess how it had changed in the absence of
161	offensive and defensive pressures. By providing the scorpions with the full length of time
162	necessary to regenerate their venom, we ensured that there was ample time for the treatments
163	to elicit a response in the chemistry of the regenerated venom.
164	
165	(d) Venom analysis
166	Venom was collected using an Arthur H. Thomas Co. Z789 Square Wave Stimulator to
167	electrostimulate the telson at approximately 25 volts (5.5 pulses/sec, for 15 milliseconds per
168	pulse). Extracted venoms were diluted in 150 μ L of degassed phosphate buffered solution
169	(PBS- Life Technologies), centrifuged for 10 minutes total at 32,000 RPM, and filtered
170	through a syringe-driven 4mm 0.22 μ m filter (Millipore). Venom profiles were obtained
171	using size-exclusion fast protein liquid chromatography (FPLC) using a Superdex [™] 75
172	10/300 (Tricorn) GL Column (13 μ m, 10mm×300mm – GE Healthcare) at 4 °C with 100%
173	PBS buffer at 0.50 ml/minute with 0.5 ml elutions for 45 mL on an ÄKTA™ FPLC (GE

Healthcare). Venom components were detected by absorbance measured at a wavelength of 280 nm. Finally, using venom collected from the same scorpions, but three weeks after the cession of the experimental treatments, we evaluated the toxicity of each venom fraction by performing toxicity assays on a human cardiac cell line to test for vertebrate toxicity (see section f), and by performing behavioural assays on crickets to test for effects on temporary or permanent invertebrate paralysis (see section g).

180

181 *(e) Statistical analysis for profile changes*

182 To compare between the venom profiles from each treatment, we split the FPLC venom 183 profile into 14 different 'fractions', and differences in the amounts of each relative to the 184 other treatments could then be evaluated statistically. To identify the different fractions, we 185 first standardised each chromatogram to an area under the curve of 1, obtained the mean 186 chromatogram for each of the four treatments by averaging all the curves within each 187 treatment. We next fitted a spline curve to each of these mean chromatograms using the 188 smooth.spline function in R [33], with the smoothing parameter, $\lambda = 0.5$ [33, 34]. The local 189 minima in these splines were then designated as boundaries between two fractions. Local 190 minima within 1 ml from each other were averaged to create a single break between fractions, 191 with one exception: the local minima values of 38.64, 38.99, 39.78, 40.51, and 40.57 ml were 192 divided into the two groups: 38.64, and 38.99; and 39.78, 40.51, and 40.57 ml, for which 193 each was averaged to describe the combined fraction separation point. Principal component 194 analysis (PCA) was then used to describe these 14 fractions across the data set [34]. 195 MANOVA and two separate, follow-up ANOVA analyses were performed to evaluate 196 treatment effects on the first two principal components. These analyses were conducted on 197 venom samples collected at three time points: prior to the initiation of treatments (t = 0 days),

at the cessation of the treatments (t = 49 days), and 21 days after the cessation of treatments (t = 70 days).

200	To evaluate the effects of predator and prey main effects on particular peaks, we
201	calculated the mean and 95% confidence intervals for each peak evaluating the difference
202	between the predator-no predator treatment means, or the live prey – dead prey treatment
203	means. The 95% confidence intervals were calculated using non-parametric bootstrapping
204	with 10,000 simulations. For each simulation, we resampled with replacement the
205	absorbances for a particular treatment (e.g., with or without simulated predator exposure) and
206	chemical fraction. Chemical fractions five through eight were not easily distinguishable and
207	likely represent a number of compounds, and therefore, we also calculated the mean
208	treatment effects and 95% confidence intervals for the sum of these fractions (summed
209	individually for each scorpion). Treatment effects were considered significant for $\alpha = 0.05$
210	when 95% confidence intervals did not overlap zero.

- 211
- 212

213 (f) Predator cell assays

214 The biological consequences of observed changes to the venom profiles were evaluated using 215 toxicity assays. Fraction concentrations were determined using the A280 method [35]. A 216 human cardiac cell line (Sciencell) was used as a vertebrate assay, following Schneider [29]. 217 Vertebrate cells were maintained and assays were performed as previously described by 218 Andreosso, Smout [36] and Chaousis, Smout [37]. An xCELLigence SP RTCA system 219 (ACEA Biosciences) with an E-plate seeded with 150 µL cardiac media (Sciencell) and 5000 220 human cardiac cells were incubated overnight at 37 °C and 5 % CO₂. 221 The cell response to each fraction (20 μ L) and 100% PBS solution (control) was measured by the xCELLigence system as changes to cell index. Cell response is a 222

combination measure of changes in media conductivity or cell contact/toxicity, which varies
as the cells deform in response to exposure to a chemical sample. The cell index readouts
were blanked against the PBS control, and the maximum drop value in 2 hours after venom
addition was deemed the predator cell response. The relative response to whole venom as a
percentage was then used to graph the activity level of the venom peaks. We used two-tailed
t-tests to compare the response of each venom fraction to the PBS control to identify peaks
that significantly altered media conductivity or cell contact/toxicity.

230

231 (g) Prey toxicity assays

232 Acheta domesticus cricket assays were performed by evaluating whether a given venom 233 fraction was active towards immature crickets. To evaluate the effects of each fraction, 3 µL 234 of one of the 14 chemical fractions was injected ventrally into the pronotum of an immature 235 cricket varying in mass from 0.1 to 0.2 grams. Immediately after injection, the cricket was 236 inserted into a clean, 9-dram clear styrene tube with snap-on lid and rolled onto its back every 237 10 seconds 18 times for a total of 3 minutes. A compromised righting response was recorded 238 when a cricket was unable to right itself within 60 seconds of being rolled onto its dorsal side. 239 Each fraction was replicated with 10 crickets. We used a 2 by 2 Fisher's exact test to compare 240 the cricket response from each venom fraction to a PBS control.

241

242 *3 Results*

243 (a) Effects of predator-prey interactions on venom composition

The experimental results were used to evaluate our hypotheses on venom plasticity using a model animal the rainforest scorpion *Liocheles waigiensis*. The venom profiles obtained from venom extraction before the experimental treatments began were not significantly different from each other (Figure 1A, see Electronic Supplementary Material 1 for statistical analysis).

248	After treatment there was a difference between the venom profiles of the predator-treated and
249	the predator-excluded scorpions. These profiles varied greatly in the relative concentration of
250	multiple chemical fractions, with the greatest difference in treatments occurring in peak
251	fraction 12 (Figure 1B). From the principal component analysis we obtained two major
252	principal components, PC1 and PC2, which explained 53.6% and 25.4% of the overall
253	variation, respectively. Venom profiles obtained from scorpions that were and were not
254	subjected to the defensive pressure treatment were found to be significantly different using a
255	MANOVA to evaluate the treatments on the principal component weightings (Table 1), with
256	increased predator exposure leading to lower and higher values of PC1 and 2, respectively
257	(Figure 2). This was most clearly associated with changes in fraction 12, which was reduced
258	in the scorpions exposed to simulated predators (Figure 1B, 2), which significantly decreased
259	with predator exposure (Figure 3). Fraction 2 also varied strongly in both principal
260	components, but not in a way that was interpretable with the experimental treatments (Figure
261	2). There were no interaction effects, nor any significant effects from prey manipulation
262	treatment (live versus dead prey) on the venom profile principal components 1 and 2 (Table
263	1). Profiles obtained after a 21-day recovery period following cession of treatments exhibited
264	similar patterns of difference between treatments (Figure 1B, C, ESM 1), but the magnitude
265	difference was reduced.

266

267 *(b) Toxicity assays*

Higher activity towards mammalian cells (>60%) was generally found in the toxin fractions containing larger proteins/peptides (fractions 2,3,4,5,7,and 8, Figure 3, ESM 1), which were likely 3-25 kDa due to the SuperdexTM 75 resin that was used [37]. One section of the profile contained many fractions (5-8) that were not easily distinguishable, significantly increased in response to simulated predator exposure (95% bootstrap confidence limits: 0.004, 0.083), as 273 did fractions 7 and 8 when evaluated individually (Figure 3). In addition, fractions 10 and 11 274 had some activity against mammalian cells, although the magnitude of these effects were 275 much lower than for other fractions (Figure 3, ESM 1). Toxicity towards crickets was 276 generally found in a fraction containing larger proteins/peptides (fractions 3-4), and the 277 fractions containing smaller compounds (fractions 8-14) (Figure 3A, ESM 1). Components of 278 each of these sections were reduced in response to simulated predator exposure (Figure 3). It 279 should be noted that many small molecules are not detectable at 280 nm and other detection 280 methods may be required. Undetected compounds were the likely source of activity against 281 crickets (>90%) exhibited by fractions 13 and 14, as the absorbance trace showed very 282 minimal contents. Example cell responses are provided in ESM 2. 283

284 (c) Comparing treatment effects with toxicity assays

285 Simulated predator exposure had the strongest effect on reducing the relative 286 production of fraction 12 that demonstrated activity on crickets, and to a lesser extent, 287 reduced the relative production of a section (fractions 3-4) that exhibited effects on both 288 crickets and mammalian cells (Figure 3). Fractions 5-8, which were not easily distinguishable 289 (Figure 1), significantly increased in response to simulated predator exposure (predator 290 treatment effect 95% limits: 0.003670351, 0.082905484) and portions of this section of the 291 profile exhibited activity on mammalian cells (Figure 3B). Fractions 1 and 11 slightly 292 increased in the presence of predators (Figure 3C), but neither of these had strong effects on 293 invertebrates or mammals (Figure 3A,B). The presentation of live versus dead prey had little 294 effect on the relative production of each chemical fraction, although it did slightly increase 295 the production of fraction 3 (ESM 1) that effected both crickets and mammalian cells (Figure 296 3A,B).

297

298 4 Discussion

320

299 Given current theory relating selection pressures to plastic changes in defence and 300 reproductive investment (e.g., Peacor, Peckarsky [38]), venomous mesopredators should shift 301 the balance of venom composition towards the defensive components when predator 302 exposure increases. In line with these predictions, we found evidence for a plastic change in 303 venom composition in response to increased perceived predation risk, showing for the first 304 time to our knowledge that organismal venom chemistry can change in response to a threat. 305 These changes imply a rerouting of resource expenditure, which may be nutritional or 306 energetic [12], to increase relative production of other venom fractions which are responsible 307 for toxicity to vertebrates. Overall, simulated predator exposure appeared to decrease relative 308 production of strong invertebrate toxins, while generally increasing the production of a 309 section of the venom profile with activity towards mammalian cells. These results suggest for 310 the first time to our knowledge that venoms can serve as inducible defences used against 311 predators. Inducible defence theory suggests that plastic defences are more likely to evolve in 312 highly variable or cyclic environments, where the fitness benefits of flexibility outweigh the 313 costs of maintaining this capacity for variability [2, 13] Venomous animals evolve vast, 314 complex armouries of peptides and proteins in their venoms [12], and it would appear that L. 315 *waigiensis* is able to modify the production of a subset of their complex venom cocktail to 316 suit a changing environment. The magnitude of the pressure to minimize venom cost and the 317 predatory pressure may also relate to how closely venom production tracks the rate of 318 ecological dynamics [17]. 319 Resource type did not elicit a response in venom chemistry. This lack of effect, which

insufficient variation in resource type, b) a time-lag in the scorpions' response which as a

was probed through removal of the need for venom expenditure, may have been due to a)

result was not detected, or c) an absence of inducible offence. It is well-documented that the

323 magnitude of environmental variability can influence both the magnitude and the speed of a 324 plastic response [39]. For example, moderate levels of herbivore damage may only induce a 325 'primed' state in plants rather than the immediate chemical response to high damage, while 326 low levels of damage may fail to provoke a plastic response at all [2]. In our experiment, we 327 introduced two resource 'types' (live or dead) to represent variation in the need for venom 328 during prey-capture. However, if the live resource type was not sufficiently different to the 329 dead resource type (i.e. often not requiring venom-use to obtain a meal) than any inducible 330 offence in the venom profile may not have been provoked. We tried to account for this by 331 feeding larger prey (i.e. larger in size than a scorpion's chelae) to encourage the need for 332 envenomation following van der Meijden, Coelho [32], but scorpions were still occasionally 333 observed to be killing their prey without stinging. Secondly, there may have been a time-lag 334 in any potential response to the treatment. In plants, induced chemical defences can be 335 mounted in response to attack, followed by a substantially longer 'relaxation' period before 336 returning to a ground state. For example, *Trifolium repens* mounts a systemic chemical 337 defence within 51 hours of herbivory, but requires at least 28 days to relax (Gomez et al. 338 2010). Similarly, an induced offense in response to prey-type may exhibit a relaxation period. 339 For example, in snails from the genus *Lacuna*, the longer an individual fed on previous diet, 340 the slower its induced morphological offense switched to a new food source [24]. Finally, 341 there may be no plastic response to variation in resource-type. This may be due to either 342 insufficient variation in prey-type in the wild to drive the evolution of a plastic response 343 capability, resembling the conditions under which constitutive defences are favoured by 344 plants, or due to sufficiently high fluctuations in prey-type to favour a bet-hedging strategy 345 rather than plasticity [40]. When traits respond to a selective pressure, evolution balances this 346 response between optimising the trait for the maximum fitness benefit and over-investing in 347 the trait to compensate for the effect of environmental stochasticity [15, 41]. Such bet

348	hedging strategies are ubiquitous in arms races [42], and may also be seen in venom-users
349	[43]. In the absence of an alternate prey which doesn't require stinging to be subdued, it may
350	be favourable to delay a plastic response (or exhibit none at all) and continue producing
351	costly venom even in the absence of live prey to ensure success in future opportunities to
352	catch a meal. Future work may be able to distinguish between these competing explanations
353	by investigating the variability of food resources in the natural habitat of L. waigiensis.
354	Induced plastic defences can stabilize populations against fluctuating predatory
355	pressures [17], and as such in ecological communities where venomous animals provide an
356	important food resource (e.g. [44]) induced defences could act as an important stabilizing
357	force for the community and diminish trophic cascades in food webs. Adaptive plasticity can
358	mitigate the effects of sudden disturbances by allowing populations to evolve sufficiently
359	quickly to survive abrupt change [45]. Phenotypic plasticity permits more time for
360	evolutionary adaptation to occur, and may reduce the degree of evolutionary change
361	necessary to track a moving optimal trade-off between the costs and benefits of venom
362	production [45]. Indeed, populations which exhibit greater phenotypic plasticity are generally
363	able to evolve more under global change and thereby adapt to changing environments [46].
364	However, plasticity may also slow the rate of evolutionary pressure by reducing the selection
365	pressure for genetic change [16]; whether or not venom plasticity should facilitate or inhibit
366	adaptation by venom-users to modified predator-prey interactions driven by environmental
367	change remains an open question.
368	In our bioassays, we found some venom fractions (e.g., fraction 3) have activity
369	against both the scorpion's cricket prey and mammalian heart cells, suggesting they may
370	serve to improve both prey capture and defence against predators. This may lead to
371	complicated tritrophic interactions where phenotypic changes in response to one arms race
372	(e.g., with predators) can alter the investment in another arms race (e.g., with prey) (Gangur

373 et al. in review). Furthermore, we identified fractions of the venom profile (e.g. fraction 1) 374 that increased in response to simulated predator exposure, but in isolation did not demonstrate 375 activity against mammalian heart cells. These fractions may have effectiveness against 376 another vertebrate biological pathway (e.g., pain activation), may interact with other fractions 377 to improve potency, or may be increased incidentally due to physiological constraints in 378 venom production. Thus, further research to clarify the role of these venom fractions in 379 predator defence may shed light on adaptive advantage of the observed phenotypic changes 380 in response to simulated predator exposure. Furthermore, we have only evaluated the effects 381 of each venom fraction on two distantly related taxa (mammalian cells and arthropods). In 382 some cases, organisms can target phenotypic changes in defence to the specific threats e.g. 383 [47, 48]. Further research exposing these scorpions to a range of predator species and 384 evaluating changes in venom composition may elucidate the specificity of this phenotypic 385 plasticity.

386

387 Venom research has historically been intently focused on human toxicity, due to obvious 388 reasons, the prevention of mortalities and a strong interest in medical advancements [49]. The 389 ecological and evolutionary perspectives that have been increasingly explored, particularly in 390 the last decade, offer critical insights into venomous animal ecology that has improved health 391 outcomes as well as enriched our understanding of venom-use and production. Indeed, if 392 plastic responses are widespread in venomous animals, antivenom production may be 393 improved by accounting for this potential source of variation by ensuring live prey or 394 simulated predation [50]. Furthermore, the potential role of venom in stabilising ecological 395 dynamics needs to be further explored as in some cases this may be a substantial factor 396 controlling community structure.

397

- **398** Data Accessibility: Venom extraction and assay data: Dryad doi:10.5061/dryad.sq2g4 [51].
- 399 Additional statistical analyses supporting this article have been uploaded as part of the
- 400 *supplementary material.*
- 401 *Ethics: There were no live vertebrate animals used as experimental subjects in this study.*
- 402 Authors' Contributions: Experiments were designed by ANG, TDN, and MJL, JES, and conducted by
- 403 ANG. Chemical analyses were conducted by ANG, MS, and DW. Mammalian and invertebrate
- 404 bioassays were conducted by MS and TDN, respectively. ANG and TDN conducted statistical
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524 **TABLES**

525

526 **Table 1:** MANOVA results from venom collected one week after the cessation of the

527 experimental treatments, which demonstrated a significant overall difference between the

528 fraction loadings from scorpions that were and were not subjected to the defensive pressure

529 treatment. There was no significant interaction effect.

Source	d.f.	Pillai	F	Df den	Р
Offence	1	0.040	0.978	2, 47	0.383
Defence	1	0.242	7.517	2, 47	0.001
Defence × Offence	1	0.036	0.887	2, 47	0.419
Residuals	48				

530

531 Table 2: ANOVA results from venom collected one week after cessation of experiments,

532 demonstrating significantly different fraction loadings between the scorpions that were and

were not subjected to the defensive pressure treatment along both PC1 and PC2. There were

- 534 no significant interaction effects.
- 535

Source	d.f.	M Sq	F	Р	
PC 1					
Prey	1	0.012	1.933	0.171	
Predator	1	0.048	7.643	0.008	
Predator $ imes$ Prey	1	0.001	0.170	0.682	
Residuals	48	0.006			
PC 2					
Prey	1	0.001	0.173	0.679	
Predator	1	0.018	5.928	0.019	
Predator \times Prey	1	0.005	1.744	0.193	
Residuals	48	0.003			

536

537

538 FIGURE LEGENDS

539 Figure 1. Averaged relative venom profiles for each of the four treatments taken: prior to treatments (A, t = 0 days), after the treatments ended (B, t = 49 days), and again after a 540 541 recovery period (C, t = 70 days) divided into 14 fractions. Relative absorbance measures the 542 absorbance units at 280 nm of any point along the venom profile relative to the point of 543 maximum absorbance in the profile. Venom profiles obtained from scorpions subjected to the 544 pressure for defensive venom (mouse exposure) treatment are given in dark blue (+ pressure 545 for offensive venom, live cricket prey) and light blue (- pressure for offensive venom, dead 546 cricket prey); profiles obtained from scorpions not subjected to this treatment are given in red (+ pressure for offensive venom) and orange (- pressure for offensive venom). 547 548 Figure 2. Loadings for principal components 1 and 2, separated into loadings for each 549 scorpion (A) and loadings for each venom fraction (B, 14 total peaks) with fraction number 550 indicated at the tips of each arrow. For clarity and consistency, colours are as per the curves 551 presented in Figure 1; pressure for defensive venom treatment is indicated in dark (+ pressure 552 for offensive venom) and light (- offensive pressure for offensive venom) blue, while data 553 from scorpions not subjected to the pressure for defensive venom treatment are in red (+ 554 pressure for offensive venom) and orange (- pressure for offensive venom). PC1 and PC2 555 described 53.6% and 25.4% of the overall variation, respectively. 556 Figure 3. Invertebrate (A) and vertebrate (B) toxicity assay results. Invertebrate toxicity was 557 measured by evaluating the proportion of crickets (10 crickets per treatment) that were 558 paralysed for longer than 60 seconds. Statistically significant difference from the control was 559 evaluated using a Fisher's Exact Test for each peak (ESM 1). Vertebrate toxicity was 560 evaluated by measuring vertebrate cell response to venom fractions relative to whole venom 561 response using the xCELLigence platform. Due to small sample volume, it was not possible

562	to completely separate fraction 5 from fraction 6. Statistically significant difference from the
563	PBS control (blanked at 0) was evaluated using a two-tailed t-test for each peak (ESM 1).
564	Panel C presents mean (and 95% non-parametric bootstrap confidence intervals) for the
565	difference between the treatments with and without simulated predator exposure. Confidence
566	intervals entirely above (or below) zero suggest significant effects of increased (or decreased)
567	production after simulated predator exposure. Differences were calculated after the
568	treatments ended ($t = 49$ days). Asterisks represent confidence intervals that do not overlap
569	zero. Chemical fractions five through eight were not easily distinguishable, and likely
570	represent multiple similarly sized compounds. Therefore, we have also calculated this
571	confidence interval separately (predator treatment effect 95% limits: 0.004, 0.083). Panel D
572	presents the relationship between invertebrate and vertebrate toxicity for each peak. The
573	feature scaling function $x' = \frac{x - \min(x)}{\max(x) - \min(x)}$ was used to convert max drop value into a
574	normalized vertebrate assay score in the range [0,1] for ease of comparison (invertebrate
575	assays were already scored in this range as a proportion of crickets out of 10 replicates
576	experiencing paralysis for >60 seconds after toxin injection). Error bars indicate standard
577	error. Due to low yield volume, the vertebrate assay score for fraction 5 includes both
578	fractions 5 and 6. Asterisks indicate significance for $alpha = 0.05$.