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**Behavioural Evidence for Polychromatic Ultraviolet
Sensitivity in Mantis Shrimp**

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1 **Behavioural Evidence for Polychromatic Ultraviolet Sensitivity in Mantis Shrimp**

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6

7 **POPULAR SUMMARY**

8 Mantis shrimp have spectacularly sophisticated eyes, with a number of unique elaborations
9 that stretch their visual capabilities far beyond our own, including deep into the ultraviolet (UV)
10 range. Bok *et al.*, 2018 uses trained and innate behavioural response experiments to show that mantis
11 shrimp are able to detect and discriminate various UV stimuli. Most notably, they respond
12 differentially to stimuli in the near-UVB (< 315 nm in wavelength) versus longer-wavelength UVA
13 stimuli. These UVB cues lie outside the discriminable range of most other animals and could afford
14 the mantis shrimp yet another covert visual signalling domain.

15

16 **KEYWORDS:** Ultraviolet vision, Visual ecology, Mantis shrimp, Colour vision

17

18 **ABSTRACT**

19 Stomatopod crustaceans are renowned for their elaborate visual systems. Their eyes contain a plethora
20 of photoreceptors specialized for chromatic and polarization detection, including several that are sensitive to
21 varying wavelength ranges and angles of polarization within the ultraviolet (UV) range (< 400 nm).
22 Behavioural experiments have previously suggested that UV photoreception plays a role in stomatopod
23 communication, but these experiments have only manipulated the entire UV range. Here, using a behavioural
24 approach, we examine UV vision in the stomatopod *Haptosquilla trispinosa*. Using binary trained choice
25 assays as well as innate burrow choice experiments, we assessed the ability of *H. trispinosa* to detect and
26 respond to narrow-band LED stimuli peaking near 314 nm (UVB) versus 379 nm (UVA) in wavelength. We
27 find that *H. trispinosa* can discriminate these stimuli, and appears to display an aversive reaction to UVB
28 light, suggesting segregated behavioural responses to stimuli within the UV range. Furthermore, we find that

29 *H. trispinosa* can discriminate stimuli peaking near 379 nm versus 351 nm in wavelength, suggesting that
30 their wavelength discrimination in the UV is comparable to their performance in the human-visible range.

31

32 **BACKGROUND**

33 Stomatopods, or mantis shrimp, are well known for their aggressive predatory behaviour,
34 sophisticated social interactions, and colourful markings (**Figure 1A**) [1]. However, they have drawn the most
35 extensive scientific interest for their unusual and complex visual systems. Their eyes are modified from
36 typical malacostracan dichromatic, apposition compound eyes by a midband of specialized ommatidia that
37 horizontally bisects each eye (**Figure 1B**). Within this midband region, the photoreceptors are structurally and
38 physiologically adapted for the detection and discrimination of eight colour bands within the human-visible
39 range (400-700 nm), as well as linearly- and circularly-polarized light [2-5]. Furthermore, the midband
40 contains up to five types of ultraviolet (UV) photoreceptors, maximally sensitive to various wavelength ranges
41 of light below 400 nm [6-8], including a pair of UV-linear-polarization-sensitive photoreceptors [9]. These
42 UV photoreceptors are uniquely tuned to narrow wavelength ranges of UV light by filtering pigments in the
43 optical elements of the ommatidia derived from biological sunscreen compounds [10-12], suggesting the
44 potential for chromatic discrimination in the UV range.

45 Despite a robust understanding of stomatopod retinal physiology, little is known about how visual
46 information is processed and employed to initiate or mediate behavioural responses. Behavioural trained
47 choice foraging experiments have been performed at human-visible wavelengths in order to demonstrate
48 sensitivity to colour [13, 14], linear polarization [15, 16], and circular polarization [5]. Furthermore, results of
49 burrow-choice experiments have suggested that sensitivity to circularly polarized light may play a role in
50 intraspecific communication [17]. Behavioural assays that test the UV range (< 400 nm) are thus far limited to
51 antagonistic encounter experiments that suggest stomatopods assess UV cues in territorial contests [18].
52 However, these experiments manipulated the entire UV range and did not examine the contribution of
53 individual UV receptor spectral types to stomatopod behavioural responses. It is not known whether the
54 multiple UV photoreceptor classes found in the stomatopod eye are used to make spectral discriminations.

55 The stomatopod *Haptosquilla trispinosa* has at least three spectral classes of UV-sensitive
56 photoreceptors located in the eighth reticular cells (R8s) of the midband (**Figure 1C-D**) [14]. Two of these
57 photoreceptor classes have segregated spectral sensitivity curves, with one photoreceptor class responding to

58 light primarily in the UVA range (315-400 nm), and a second absorbing light strongly in the UVB range (\leq
59 315 nm). Based upon these electrophysiological spectral sensitivity measurements, we hypothesized that *H.*
60 *trispinosa* would be capable of detecting and discriminating UVA versus UVB stimuli. Here we present the
61 results of trained predatory choice and innate burrow preference behavioural experiments in *H. trispinosa*. We
62 show that these stomatopods are able to discriminate and behaviourally respond to UVA and near-UVB
63 stimuli. Furthermore, our results suggest spectrally distinct roles for UVA versus UVB cues in stomatopod
64 behaviour, possibly related to intraspecific communication.

65

66 MATERIALS AND METHODS

67 *Animals*

68 *Haptosquilla trispinosa* individuals were collected at the Lizard Island Research Station (Queensland,
69 Australia, 14°40'43.9"S, 145°26'47.9"E) at 1 meter depth in May and June of 2012 (for trained choice tests)
70 and in June 2014 (for the innate burrow preference tests). The individuals used in the trained choice
71 experiments ranged in length from 23-36 mm, with a mean of 28.9 mm. The individuals used in the innate
72 burrow choice experiments ranged in length from 21-37 mm, with a mean of 28.7 mm. They were kept for 1-5
73 days in individual cups with daily water changes and regular feeding with small pieces of snail or crustacean
74 meat until they were moved to the training or experimental setups.

75

76 *Trained Choice Tests*

77 The trained choice test followed a similar approach to that of Thoen et al. [14]. The animals were
78 housed in individual aquaria with artificial burrows constructed from plastic vials positioned in a sand bed and
79 with constant seawater flow-through. The training and experimental apparatus was custom constructed by
80 John Cataldi at the University of Maryland, Baltimore County MME Technical Service Center Machine Shop.
81 It consisted of pair of submersible targets with a 3 mm hole in the centre connected to an above-water light
82 emitting diode (LED) mount and controller by a pair of 10-cm-long, 3 mm diameter optical guides in
83 blackened brass tubes (**Figure 1E, S1**). The stimuli were generated using one of three LEDs with maximum
84 emission (λ_{max}) at 314.3 nm (UVB), 378.3 nm (UVA) and 351.1 nm (UVA-351) (**Figure 1E, inset**). Note that
85 UVA refers to the 378.3 nm LED stimulus unless otherwise indicated. Wavelengths of light beyond 400 nm
86 were blocked by a UV bandpass filter in the target head. Refer to **Figure S1** for diagrammatic representations

87 and technical details of the testing apparatus and the stimuli. The brightness of the LEDs were modulated by a
88 custom controller (**Figure S1E, S2A-B**).

89 The trained choice test exploited an innate predatory behaviour where *H. trispinosa* will lunge from
90 its burrow to attack a target. In the choice test trials, the animals were presented with a pair of targets and
91 trained to associate a food reward (shrimp or mollusc meat) with a UVA or UVB stimulus. Possible
92 confounds in brightness were controlled for by modulating the relative intensities of the two LED stimuli in
93 the trials. The intensity of each LED was randomized between five intensity settings in the UVA vs. UVB
94 experiments (Figure S2A) and between four settings in the UVA vs. UVA-351 experiment (Figure S2B).

95 Prior to the trials, individuals were first made accustomed to feeding from a single practice target,
96 presented alone and without any emitted UV stimuli. The underside of these training targets had a small
97 groove, not visible from the front, that a small piece of snail meat could be affixed to. The choice trial targets
98 used in the subsequent experimental rounds did not have this groove and never came in contact with food.
99 Individuals that learned to feed from the single training target were then split into cohorts of initially 12
100 randomized individuals, roughly balanced for even distributions of gender and size, to begin choice training.
101 The animals in each cohort were trained to associate a particular stimulus with a reward in a randomized
102 binary choice context against a second alternate stimulus. For training, a piece of food was affixed to the
103 underside of the correct training target. The two targets were positioned in the water, at a distance from the
104 burrow necessitating the animal to lunge fully from the burrow in order to collect the food. Once the animals
105 were feeding from the correct stimulus consistently, choice test trials were initiated.

106 Trials were carried out by first blocking the burrow entrance with an opaque plastic sheet so that
107 enthusiastic individuals were prevented from leaving the burrow until the targets were positioned properly.
108 Once the targets were in place, seawater that had contained thawing reward food (shrimp or snail muscle) was
109 poured broadly over the front of the sheet to alert and stimulate the animals with an odorant cue. The plastic
110 sheet was then lifted, and the animals were given two minutes to make a choice. Choices were scored when an
111 animal fully exited the burrow and touched one of the targets. Correct choices were rewarded by giving the
112 animal a piece of food on a feeding stick. Incorrect choices terminated the trial for that individual and the
113 targets were removed. If no choice was made, it was noted whether the animal extended its head from the
114 burrow to assess the target but never attacked, or did not emerge at all (**Figure S1B**). The experimenter
115 observed the trials on a camera viewscreen from behind masking material mounted on the LED controller.

116 Ambiguous responses were not rewarded, but were reviewed and scored from the recorded video. If the
117 animal did not clearly hit a target, or if it lunged between the targets, it was scored as a failure to participate.

118 Trials were carried out three to five times per day until each cohort reached 30 trials. Intermittent
119 binary training rounds were performed (as described above) once or twice a day, usually before the first trial
120 round of the day, in order to motivate and reinforce the behaviour. The cohort that was trained to choose UVA
121 from UVB was used for control experiments where they were presented with two identical UVA stimuli. This
122 same cohort was then used in an additional 30 trial experiment that asked the individuals to continue choosing
123 the UVA stimulus, but now against the UVA-351 stimulus. The initial experiments involving cohorts 1, 2, and
124 3 (UVA vs. UVB, UVB vs. UVA, and UVB vs. dark) were carried out simultaneously. A second set of
125 experiments involving cohorts 4 and 1 (dark vs. UVB, and UVA vs. UVA-351) were performed
126 simultaneously following the initial three. During the experiments, trials were performed throughout the day
127 and we alternated from one cohort to another after each trial round.

128 Upon the completion of the trials, correct, incorrect, and non-participatory outcomes were collated for
129 each individual (Table S1). An individual did not participate (DNP) if it assessed the targets but made no
130 choice. All individuals who never made a choice in a trial were removed from the dataset. In some cohorts,
131 this created the gender ratio imbalance reported in Table S1. The percent correct choices and percent
132 participation values for each individual were then averaged within each experimental cohort in order to
133 perform statistical analysis without pseudoreplication (**Table 1**, see additional details about statistical analysis
134 below).

135

136 *Innate Burrow Preference Tests*

137 The second set of experiments exploited innate cover-seeking behaviour in *H. trispinosa*. Naive
138 animals, assorted into four groups with roughly equivalent gender and body length distributions, were
139 introduced into the centre of a circular arena facing a pair of artificial burrows that emitted either UVA ($\lambda_{\max} =$
140 379.1 nm), UVB ($\lambda_{\max} = 317.4$ nm), or no light stimuli (**Figure 1G**). Note that these stimuli differ slightly in
141 spectral properties from those in the trained choice tests despite being generated by the same LEDs because of
142 different optical components in their respective setups (**Figure S1, S2**). In the experiment that tested UVA
143 versus UVB preference, the LEDs were modulated in order to produce stimuli of equivalent sum radiance
144 (**Figure S2C**). The stimuli were again generated by a pair of LEDs which illuminated a diffuser at the back of

145 the burrow, and passed through a 1 cm hole covered by a UV bandpass filter into the burrow. The stimuli
146 were randomized between the two burrows for each trial. The arena had a sand bottom and continuous water
147 flow-through entering from behind the animals and exiting through blackened air line tubes emerging through
148 the bottoms of the two artificial burrows. Refer to **Figure S1F-H and S2C** for diagrammatic representations
149 and technical details of the testing apparatus and the stimuli.

150 For each trial, an animal was loaded into a stoppered clear glass flask and introduced into the centre of
151 the arena with the flask's opening facing the two burrow options. The stopper was removed, and the animal
152 was given up to two minutes to make a choice (it typically required much less time). Trials were observed by
153 the experimenter, positioned outside of the view of the arena, via a camera viewscreen. Choices were counted
154 when an individual moved directly from the flask into one of the burrows. In the event that the animal never
155 left the flask, never entered a burrow, or moved to an adjacent side of the arena before entering one of the
156 burrows, the trial was scored as a failure to participate. The sand at the bottom of the arena was stirred
157 between trials to obscure chemical cues.

158

159 *Statistical Analysis*

160 All statistical analyses were conducted in R (version 3.3.2 [19]). In the trained choice tests, correct,
161 incorrect, and non-participatory outcomes were recorded for each individual trial and were analysed using
162 generalized linear mixed model (GLMM) (lme4 package [20]). To assess the influence of the wavelength and
163 stimulus brightness on the choice of the individual, a single model was conducted on the binary response
164 variable using the fixed factors of wavelength, relative brightness level, and the interaction between these two
165 factors. We included the individual animal identity as a random term to control of the repeated measures per
166 animal. Single term deletions were used to reduce the model to its minimum form.

167 As brightness changes were found not to influence the choice of the animals, further analyses for
168 assessing the effect of the wavelength on choices were conducted using a one-sample Wilcoxon test. In each
169 experiment, the repeated results from the multiple trials per individual were averaged to provide a mean
170 proportion for a measure of successes for each individual (**Table S1**). These data were then compared for each
171 experiment against an expected mean of 0.5 (50% correct choices) based on a null hypothesis of the animals
172 not having the capability to differentiate between the different spectral contents of the stimuli (**Table 1**).

173 For the burrow preference tests, single naïve individuals were tested once in each trial and a binomial
174 test was used to analyse if their pooled innate responses differed due to the different burrow illumination
175 stimuli.

176

177 RESULTS

178 *Trained Choice Tests*

179 Six experiments were performed testing the ability of *H. trispinosa* to learn to discriminate and
180 choose UVA ($\lambda_{\max} = 378.3$ nm) and UVB ($\lambda_{\max} = 314.3$ nm) stimuli (**Table 1, Figure 1F**). We found that
181 while Cohort 1 was able to choose the UVA versus the UVB stimulus at a significant rate (89.5% success;
182 Wilcoxon, $V = 28$, d.f. = 1, $p = 0.022$), Cohorts 2 and 3 were unable to differentiate between the UVB
183 stimulus versus the UVA stimulus (65.1% success; Wilcoxon, $V = 25$, d.f. = 1, $p = 0.341$), or versus a dark
184 stimulus (41.3% success; Wilcoxon, $V = 10$, d.f. = 1, $p = 0.291$). Attempts to train Cohort 4 to choose a dark
185 stimulus versus a UVB stimulus were also unsuccessful (58.8% success; Wilcoxon, $V = 27$, d.f. = 1, $p =$
186 0.234). Cohort 1 was then used in a control experiment where the stomatopods were presented with an
187 identical pair of UVA stimuli. The cohort's preference for the two stimuli was identical (46.1% preference for
188 the left target; Wilcoxon, $V = 12$, d.f. = 1, $p = 0.799$). Finally, Cohort 1 was tested to again choose UVA
189 versus a UVA-351 ($\lambda_{\max} = 351.1$) stimulus, only 27.2 nm apart in maximum emission. The cohort remained
190 able to choose the UVA stimulus at a significant rate (64.1% success; Wilcoxon, $V = 35$, d.f. = 1, $p = 0.015$).
191 See **Table S1** for choice and participation data for each individual in these experiments. Over all the tests, the
192 relative brightness of the stimuli did not have any effect on the choice of stimulus (GLMM, $\chi^2 = 1.2552$, d.f. =
193 2, $p = 0.534$), however there was a clear difference in how the animals responded to the spectral pairs
194 (GLMM, $\chi^2 = 16.913$, d.f. = 2, $p < 0.001$) (**Figure S3**). The full dataset used in the GLMM analysis can be
195 found in Supplementary Data File 1, with an explanation of its contents in the Supplemental Materials
196 document.

197 In the trained choice assays we observed a trend in participation related to the UVB stimuli.
198 Participation was markedly reduced in experiments where we attempted to train the stomatopods to choose the
199 UVB stimulus (UVB vs. UVA, 22.3% participation; UVB vs. dark, 45.3% participation) (**Table 1**). A similar
200 effect was also noted when a UVB stimulus was present as an alternative target stimulus (UVA vs. UVB,
201 56.5% participation, dark vs. UVB, 46.2% participation). When there was no UVB stimulus presented in the

202 experiment, the animals displayed an elevated rate of participation (UVA vs. UVA control, 95.7%
203 participation; UVA vs. UVA-351, 69.9% participation).

204 Males and females did not differ markedly in the percent correct choices in any of these experiments
205 (**Figure 2A**). However, males invariably showed much lower participation than females when there was a
206 UVB stimulus present (**Figure 2B**) (males, 26.3% participation, n=17; females, 57.3% participation, n=17;
207 Wilcoxon, $W = 55.5$, d.f. = 1, $p < 0.001$).

208

209 *Innate Burrow Preference Tests*

210 In the innate burrow choice tests, *H. trispinosa* showed significant aversion both to UVA ($\lambda_{\max} =$
211 379.1 nm) and UVB ($\lambda_{\max} = 317.4$ nm) stimuli (**Figure 1H, Table 2**). When presented with a choice between
212 UV emitting burrows and a dark burrow, they chose the dark burrow at a significant rate: 83.3% preference
213 versus UVB (Binomial test, number of dark burrow choices = 35, number of choices = 42, $p < 0.001$), and
214 74.4% preference versus UVA (Binomial test, number of dark burrow choices = 32, number of choices = 43, p
215 < 0.001). Based on the apparently greater aversion to UVB stimuli observed in the trained choice tests, we
216 hypothesized that when given an intensity-matched choice between burrows emitting UVA versus UVB light,
217 *H. trispinosa* would prefer UVA emitting burrows. However, we found no significant preference in this case
218 (54.2% preference for UVB; Binomial test, number of UVB burrow choices = 13, number of choices = 24, $p =$
219 0.838). When both burrows were dark, *H. trispinosa* showed no significant side preference (55.0% preference
220 for the left burrow; Binomial test, number of left burrow choices = 11, number of choices = 20, $p = 0.824$).
221 We also observed depressed participation in the UVA versus UVB experiment (52.2%) compared to
222 experiments with a dark burrow option (UVA vs. dark, 80.8%; UVB vs. dark, 75.4%; and dark vs. dark,
223 74.1%).

224

225 **DISCUSSION**

226 The behavioural experiments demonstrate spectral discrimination within the UV range in the
227 stomatopod species *Haptosquilla trispinosa*, regardless of brightness cues (consistent with previous
228 experiments at human visible wavelengths showing that this species does not appear to use brightness cues
229 when making colour discriminations [14]). The results of the trained choice experiments demonstrated that *H.*
230 *trispinosa* could learn to choose a UVA stimulus against a UVB stimulus, but not *vice versa* (**Figure 1F**,

231 **Table 1**). This result could be interpreted as *H. trispinosa*'s simply being unable to detect UVB light.
232 However, the experiments also revealed a depressed participation rate when we attempted to train the
233 stomatopods to choose a UVB stimulus, or simply when a UVB stimulus was presented as an alternate target
234 choice (**Figure 1F**). This observation led us to hypothesize that *H. trispinosa* could discriminate the UVB and
235 UVA stimuli from one another, but innately treated the UVB stimulus as an aversive cue and could therefore
236 not learn to associate it with a food reward. We confirmed the ability of *H. trispinosa* to detect UVB by the
237 innate burrow choice experiments, where we found that individuals consistently chose a dark burrow over a
238 UVB-emitting burrow (**Figure 1H, Table 2**). We further hypothesized that when given the choice of a UVA-
239 versus UVB-emitting burrow, the stomatopods would prefer the UVA-emitting burrow. However, we instead
240 found that they had no preference and diminished participation in this case, suggesting that they simply do not
241 like any brightly lit burrows. Taken together, these experiments show that *H. trispinosa* can discriminate UVA
242 and UVB cues. However, they appear to treat the cues differently in the two behavioural contexts; being
243 averse to the UVB stimulus in predatory behaviours and avoiding both UVA and UVB in shelter seeking
244 behaviours. Furthermore, the depressed participation by males in the trained choice tests when the UVB
245 stimulus was present suggests a sexually dimorphic response to UVB cues (**Figure 2**).

246 UV sensitivity and UV-cue-driven behaviours are common amongst animals (reviewed in [21]), but
247 the majority of examples involve UVA photoreception. Recently, behaviourally relevant UVB sensitivity has
248 begun to receive some attention [22-24]. For instance, thrips display a UVB-specific phototactic response for
249 an unknown purpose [22], and jumping spiders use UVA and UVB reflective patches in conjunction as sexual
250 signalling cues [23, 24]. Our results offer an aquatic example of UVB behavioural sensitivity, which is
251 somewhat surprising since UVB light is rapidly attenuated in water [21]. However, many stomatopods,
252 including *H. trispinosa*, live in shallow, clear tropical waters with abundant UVB irradiance.

253 It is not well understood how stomatopods process colour information, or whether UV photoreceptors
254 are integrated into the longer-wavelength colour processing system. The subset of photoreceptors responsible
255 for UV sensitivity (the R8s) project directly to the medulla, bypassing the lamina where the projections of the
256 R1-7 receptors (maximally sensitive to wavelengths of light between 400 and 700 nm) terminate, and where
257 spectral comparison is thought to be initiated [25]. Interestingly, we found that stomatopods could
258 discriminate cues within the UVA range that emitted light maximally at 378.3 nm (UVA) and 351.1 nm
259 (UVA-351), only 27.2 nm apart in maximum emission (**Figure 1F, Table 1**). This spectral discrimination

260 performance is comparable to the capabilities of this species at human-visible wavelengths [14]. Since only
261 the midband row 1 R8 absorbs strongly in this region, it is likely that this discrimination is facilitated in
262 conjunction with the third R8 receptor (maximally sensitive around 325 nm, **Figure 1C**, dashed grey line) or
263 the R1-7 receptors. One of these R1-7 receptors, the midband row 4 distal main rhabdom receptor, is
264 maximally sensitive at 420 nm but overlaps significantly in sensitivity with the row 1 R8, down to around 370
265 nm [8]. This suggests the potential for spectral comparison between the R8s and main rhabdom receptors in
266 the midband.

267 Recent wavelength discrimination experiments in *H. trispinosa* at human-visible wavelengths (400-
268 700 nm) have implied that stomatopods may be using a novel form of chromatic processing, recognizing
269 narrow bins of wavelengths in a manner that can be likened to a spectral barcode scanner [14]. Such a system
270 could rapidly encode and assess specific colour patterns, such as the resplendent markings found on many
271 species of mantis shrimp. It remains to be seen how the UV-sensitive photoreceptors would contribute to such
272 a chromatic processing system. However, our results demonstrating UV discrimination and UVB-specific
273 aversion raises the exciting potential for the presence of UVB-encoded aggression cues on stomatopods that
274 could serve as a robust and covert means of identifying one another's intentions before coming to blows.

275

276 **ETHICS STATEMENT**

277 Research using *Haptosquilla trispinosa* was carried out under the supervision of the staff of Lizard
278 Island Research Station and with the following requisite permits: Australian Marine Parks (GBRMPA) permit
279 nos. G12/35005.1, G14/36625.1 and Fisheries Act no. 140763.

280 **DATA ACCESSIBILITY STATEMENT**

281 The datasets supporting this article have been uploaded as part of the supplementary material.

282 **COMPETING INTERESTS**

283 We have no competing interests.

284

285 **AUTHOR CONTRIBUTIONS**

286 MJB designed the project, carried out the experiments, analyzed the data, and edited the manuscript. NWR
287 analyzed the data and edited the manuscript. TWC assisted in experiments, supervised the project, and edited
288 the manuscript.

289

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303

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364

365 TABLES

366

367 **Table 1. *Haptosquilla trispinosa* trained choice results.** The “exp. stimuli” column indicates the wavelength
 368 that each cohort was trained to choose listed first. UVA, $\lambda_{\max} = 378.3$ nm; UVB, $\lambda_{\max} = 314.3$ nm; UVA-351,
 369 $\lambda_{\max} = 351.1$ nm. N , number of individuals in each cohort. Percent participation (% part.) and percent correct
 370 choices (% cor.) are calculated by averaging the percent participation and percent correct choices for each
 371 individual in each experiment (see **Table S1** for full choice results for all individuals). P-values are calculated
 372 using percent correct choices for each individual within each experiment in a Wilcoxon signed rank test with
 373 continuity correction ($P_0 = 0.5$, $\alpha = 0.05$).

<u>exp. stimuli</u>	<u>cohort</u>	<u>N</u>	<u>% part.</u>	<u>% cor.</u>	<u>Wilcoxon p-value</u>	
UVA vs. UVB	1	7	56.5	89.5	0.0215	*
UVB vs. UVA	2	9	22.3	65.1	0.3411	<i>ns</i>
UVB vs. dark	3	8	45.3	41.3	0.2912	<i>ns</i>
dark vs. UVB	4	10	46.2	58.8	0.2340	<i>ns</i>
UVA vs. UVA ^a	1	7	95.7	46.1 (L)	0.7988	<i>ns</i>
UVA vs. UVA-351	1	8	69.9	64.1	0.0156	*

374 ^aIn control experiments the same cue is present at both targets. “L” or “R” denote if the animal chose the left or right option.

375

376 **Table 2. *Haptosquilla trispinosa* burrow preference experimental results.** UVA, $\lambda_{\max} = 379.1$ nm; UVB,
 377 $\lambda_{\max} = 317.4$ nm; P-values are calculated with a binomial test ($P_0 = 0.5$, $\alpha = 0.05$).

<u>exp. stimuli</u>	<u>preferred</u>	<u>alternate</u>	<u>choices</u>	<u>DNP^b</u>	<u>trials</u>	<u>% part.</u>	<u>% pref.</u>	<u>p-value</u>	
UVB vs. dark	35 (dark)	7 (UVB)	42	10	48	80.8	83.3	1.51x10⁻⁵	*
UVA vs. dark	32 (dark)	11 (UVA)	43	14	54	75.4	74.4	1.91x10⁻³	*
UVB vs. UVA	13 (UVB)	11 (UVA)	24	22	46	52.2	54.2	0.838	<i>ns</i>
dark vs. dark ^a	11(L)	9 (R)	20	7	35	74.1	55.0	0.824	<i>ns</i>

378 ^aIn control experiments the same dark cue is present at both burrow options. “L” or “R” denote if the animal chose the left or right option.

379 ^bDid not participate: Did not directly enter one of the choice burrows within the experimental period.

380

381 **FIGURES LEGENDS**

382

383 **Figure 1. *Haptosquilla trispinosa* UV behavioural assays.**

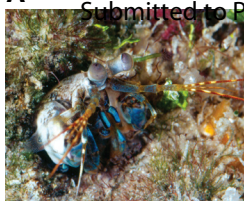
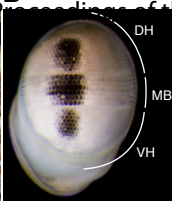
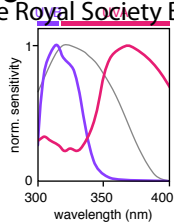
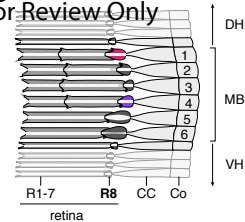
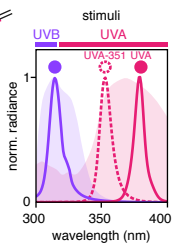
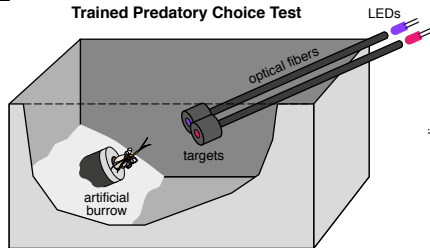
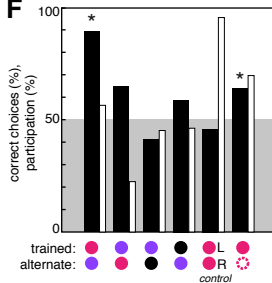
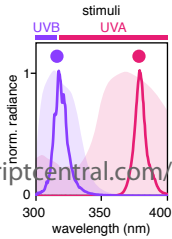
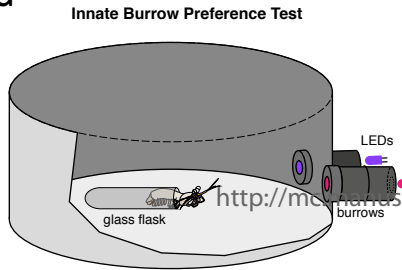
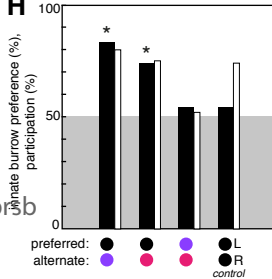
384 (A) *H. trispinosa* pictured in a natural burrow. Photo: Roy Caldwell. (B) An *H. trispinosa* eye with the dorsal
385 hemisphere (DH), ventral hemisphere (VH), and midband (MB) labelled. (C) Spectral sensitivities of the three
386 *H. trispinosa* UV sensitive photoreceptors, adapted from electrophysiological recordings in Thoen et al. [14].
387 UVA and UVB regions of the spectrum are indicated. (D) A diagrammatic cross section through the midband
388 of the eye. Ommatidial components: R1-7, reticular cells 1-7; R8s, reticular cell 8; CC, crystalline cone; Co,
389 cornea. R8 cell colour corresponds to spectral sensitivities in C, and their locations are inferred from typical
390 opsin expression patterns [10, 26] and UV filter pigment localization [11]. (E-H) Results of trained predatory
391 choice (E-F) and innate burrow preference (G-H) behavioural assays. E and G show simplified diagrams of
392 the respective experimental setups with normalized radiance spectra of their stimuli displayed to the left. Full
393 schematics and additional details can be found in **Figure S1**. Shaded colour regions correspond to R8 spectral
394 sensitivities in C. The bar graphs display average correct choice percent in the trained choice tests (F, black
395 bars) and burrow preference percent (H, black bars), as well as respective percent participation (thin, white
396 bars). Both bar graphs are labelled with circles indicating their respective stimuli corresponding to the spectral
397 plots in E and H. Black circles indicate a dark stimulus. Trained choice significance in E is calculated using a
398 Wilcoxon signed rank test with continuity correction from the percent correct choices for each individual in
399 the cohort ($P_0 = 0.5$, $\alpha = 0.05$; Table 1). Innate burrow preference significance in H is calculated with a
400 binomial test ($P_0 = 0.5$, $\alpha = 0.05$; Table 2).

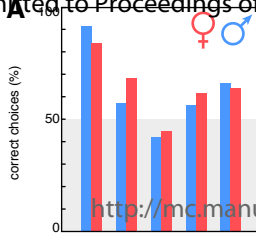
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402 **Figure 2. Male (blue bars) and female (red bars) correct choices (A) and participation (B) in trained**403 **choice experiments from Figure 1F.** Symbol circles for tests are as in Figure 1F. Percentages are derived

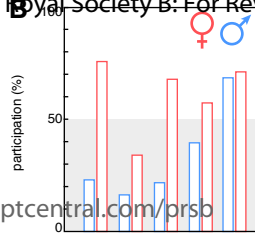
404 from pooled choice data for each gender (Table S1).

405

A**B****C****D****E****F****G****H**



trained: ● ● ● ● ●
 alternate: ● ● ● ● ●



trained: ● ● ● ● ●
 alternate: ● ● ● ● ●

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