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- 1 Do crab spiders perceive Batesian mimicry in hoverflies?
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Abstract

13	Many putative Batesian mimics only approximately resemble their supposed models, and such
14	"imperfect" mimics are readily distinguished from defended species by humans and other
15	vertebrates. One explanation for the existence of imperfect mimics is that the most important
16	predators of many mimics have very different sensory and cognitive abilities from those of a typical
17	vertebrate. In such circumstances, selection for more accurate mimicry, as perceived by humans,
18	may be reduced. Little is known, however, about how invertebrate predators perceive and respond
19	to mimicry in insect prey. Here, we investigate the foraging behaviour of the crab spider Synema
20	globosum, an important predator of flower-visiting insects at our field site which frequently
21	encounters both Batesian mimics (hoverflies - Diptera: Syrphidae) and their models (bees and wasps
22	- Hymenoptera). In the field, we found that spiders can distinguish among dipteran and
23	hymenopteran prey taxa, frequently attacking some models and mimics, but avoiding others.
24	Laboratory experiments suggest that some apparently accurate mimic taxa are more likely to be
25	avoided when spiders have prior experience of an aversive wasp model. Avoidance by spiders of
26	black and yellow striped artificial prey suggests visual cues play a role in prey selection, but there
27	was no evidence that olfactory cues are used to identify dangerous or noxious species. Overall, our
28	results provide some support for the hypothesis that invertebrate predator behaviour can generate
29	selection on visual signals in putative Batesian mimics.

#### Introduction

32 Batesian mimics are palatable and harmless species which resemble aversive or defended prey, 33 many of which advertise their unprofitability to predators with aposematic warning signals (Bates, 34 1862; Ruxton et al., 2004). They gain protection from attack by fooling predators into misidentifying 35 them as unprofitable or dangerous. While for over a century Batesian mimicry has been considered a 36 classic example of adaptive evolution, it has long been recognised that many supposed mimics only 37 approximately resemble their putative models (Gilbert, 2005; Sherratt, 2002). This poses the 38 question: if natural selection by predators favours the ever closer resemblance of a mimic to its 39 model, why are so many mimics clearly "imperfect" (Edmunds, 2000)? There are several possible 40 answers to this question (reviewed in Gilbert, 2005), with recent evidence suggesting that selection for perfect mimicry may be relaxed in palatable but relatively unprofitable prey (Penney et al., 41 42 2012), and that mimetic accuracy may be traded-off against thermoregulatory costs (Taylor et al., In 43 Review). However, there is no consensus about the most important factors influencing the precision 44 with which mimics resemble their models (Gilbert, 2005).

45 A widely recognised problem with studies of mimicry is that, whilst many mimics appear imperfect 46 to human or other vertebrate eyes, the predators that are actually responsible for selection of 47 mimetic phenotypes may include taxa (e.g. insects and spiders) with very different sensory and 48 cognitive abilities (Stevens, 2007). Thus, it is entirely possible that real-world predators perceive apparently imperfect mimics as being indistinguishable from aversive or defended prey (Cuthill and 49 50 Bennett, 1993). As yet, however, there is very little empirical evidence for differences in perception 51 of prey signals among predators, especially in natural situations, and invertebrate predators in 52 particular have been neglected in the literature on mimicry (but see Kauppinen and Mappes, 2003; 53 Rashed et al., 2005). This gap in our knowledge, combined with the fact that we know almost 54 nothing about the relative importance of different current and historical causes of selection on

warning and mimetic patterns in natural populations of invertebrates, seriously limits our ability to
test hypotheses about the evolution of aposematism and mimicry.

57 It was noted as far back as the origin of the theory of Batesian mimicry that a wide range of 58 predators avoided aposematic butterflies including dragonflies, mantids and flies (see Carpenter and 59 Ford, 1933), but to date the majority of studies of predator perceptions of aposematism and 60 Batesian mimicry have focussed on birds (e.g. Dittrich et al., 1993; Ham et al., 2006; Mostler, 1935) 61 and other vertebrates (e.g. Hetz and Slobodchikoff, 1988; Nonacs, 1985; Osorio et al., 1999). Birds 62 have been seen as important predators of a wide range of aposematic and mimetic prey taxa, as well 63 as being easily trained and having a predictable behavioural repertoire, making them ideal study species. Early experiments demonstrated that avian predators have the potential to select for 64 65 Batesian mimicry, but that they are not fooled by relatively imperfect mimics (Mostler, 1935). These 66 experiments showed that the more closely hoverflies (Diptera: Syrphidae) appeared to mimic 67 hymenopteran models, the more protection they received from predation. They also showed that 68 birds were less likely to attack certain mimics following experience with their proposed 69 hymenopteran model. Overall, these experiments suggested that birds were generally deceived by 70 mimics, but that the extent of protection enjoyed was dependent on the closeness of the mimic's 71 resemblance to its model.

Although birds demonstrate some differences from humans in the perception of prey signals (Bain et al., 2007; Dittrich et al., 1993), and may have some cognitive and sensory limitation which impact on their ability to identify prey correctly (Chittka and Osorio, 2007; Kikuchi and Pfennig, 2010), many mimetic hoverfly species appear to gain little protection from birds as a result of their resemblance to their hymenopteran models (Dlusskii, 1984). This would suggest that the potential for avian predation to select for inaccurate Batesian mimicry in invertebrate prey is limited. However, mimetic hoverflies are also attacked by many invertebrate predators (Howarth and Edmunds, 2000), and these species are likely to view the mimetic patterns of their prey in a very different way fromhumans and other vertebrates.

81 In spite of the strong evidence for the visual and cognitive abilities of invertebrates (e.g. Dyer and 82 Chittka, 2004; Pohl et al., 2008; Tibbetts, 2002), few studies have considered invertebrate predator 83 responses to warning signals and to putative Batesian mimicry. Dejean (1988) found that hunting 84 workers of the ant species, Odontomachus troglodytes, not only learned to avoid the warningly 85 coloured larvae of an African chrysomelid beetle, but also retained this behaviour for up to 28 days 86 following an initial costly experience. Similarly, mantids can learn to avoid aposematic prey 87 (Gelperin, 1968), and extend their avoidance to similarly-coloured palatable mimics (Berenbaum and 88 Miliczky, 1984), and palatable, non-mimetic flies given artificial warning signals gained protection 89 from predation by dragonflies (Kauppinen and Mappes, 2003). Taken together, these studies 90 strongly suggest that invertebrate predators have the potential to play a role in selecting for 91 aposematism and Batesian mimicry (but see Rashed et al., 2005 for evidence to the contrary). 92 Spiders are probably important predators of many aposematic and mimetic invertebrates: we know, 93 for example, that crab spiders (Thomisidae) are a major source of mortality in honeybees (Apis 94 mellifera) in some circumstances (Reader et al., 2006). However, we have no direct estimate of the 95 relative contributions of spiders and other predators (e.g. birds) to selection on invertebrate colour 96 patterns, and the extent to which spiders detect and respond to aposematism and mimicry remains 97 almost completely unexplored. Spiders appear to be capable of learning to avoid signals associated 98 with a cost (Huang et al., 2011; Vasconcellosneto and Lewinsohn, 1984), and some species have

99 been shown to have an innate aversion towards defended prey (Nelson and Jackson, 2006). Most 100 spiders are thought to have relatively poor vision, and rely mainly on other senses such as olfaction 101 and mechanoreception when foraging (Herbstein, 2011). Thus, it has been suggested that spiders 102 may select for prey warning signals and mimicry in other modalities (Vasconcellosneto and 103 Lewinsohn, 1984). However, some spiders do rely heavily on vision while hunting. Jumping spiders (Salticidae) have remarkably acute vision, which may be equivalent to that of some birds (Williams
and McIntyre, 1980). They are known to rely on visual cues for the identification of palatable prey,
predatory threats and other aspects of their environment (Harland and Jackson, 2004). Furthermore,
they have been proposed as important agents of selection on some mimetic signals; juveniles of
myrmecomorphic salticids appear to benefit from Batesian mimicry of ants by avoiding attacks from
larger salticids (Huang et al., 2011).

110 Like salticids, crab spiders are cursorial (non-web-building) predators of various aposematic and 111 mimetic invertebrates (Morse, 2007; Reader et al., 2006). Relatively little is known about crab spider 112 vision, but they are considered to be predominantly visual foragers, being able to resolve objects at 113 a resolution similar to that achieved by the compound eyes of some insects (Schmalhofer & Casey, 114 1999). Observations of the crab spider *Misumena vatia* suggest that it relies heavily on 115 mechanoreceptors for prey detection, and may attack prey relatively indiscriminately (Morse, 2007). 116 The ability of crab spiders, however, or any other spiders, to detect or respond to differences 117 between defended and undefended, or mimetic and non-mimetic prey, has not been examined in 118 detail (but see Tyshchenko, 1961). Hence, the potential of spiders to select for Batesian mimicry 119 remains unexplored, and the consequences of the differences between their sensory and cognitive 120 abilities and those of vertebrates for the evolution of mimetic accuracy in prey such as hoverflies are 121 unknown.

122 In this study, with a series of field and laboratory experiments, we explore the potential of a crab 123 spider (*Synema globosum* Franganillo) to discriminate among aposematic, defended Hymenoptera 124 and visually mimetic hoverflies. We ask whether venomous wasps and bees are aversive to spiders, 125 and whether prior experience with such species makes spiders more likely to avoid their supposed 126 Batesian mimics. Finally, we ask whether any discrimination among prey types is achieved through 127 the use of visual or olfactory cues. Our ultimate objective is to shed light on the hypothesis that

- 128 predators with relatively poor visual acuity compared with vertebrates could be responsible for the
- 129 evolution of approximate or imperfect visual mimicry in their prey.

130

#### Methods

#### 133 Study System

134 We examined behavioural responses of S. globosum to common aposematic and mimetic prey, and 135 artificial models of prey, at our field site in Sobreda de Caparica, Lisboa, Portugal (38°33'67"N, 136 009°11'34"W). Adult S. globosum are the most abundant flower-dwelling predators at our site in 137 spring (Ibarra, 2013), and are frequently observed attacking flower-visiting arthropods, including a 138 range of mimetic hoverflies, and other non-mimetic flies. Although S. globosum frequently kills 139 honeybee workers (Reader et al., 2006), we have not observed it attacking any of the venomous 140 social wasps (Hymenoptera: Vespidae) which visit flowers to feed on nectar and/or hunt for prey, 141 the most common of which is the paper wasp Polistes dominula (also referred to as P. dominulus). P. 142 dominula appears to be a threat to S. globosum: spiders were attacked and consumed by wasps 143 when kept in captivity together.

144 For our experiments, we collected individuals of seven species of Diptera and two species of 145 Hymenoptera from flowers on which S. globosum is known to forage (Figure 1; Table 1). Four of the 146 dipteran species have black and yellow or orange stripes on the abdomen, and appear to be mimics 147 of vespid wasps such as *P. dominula*, one is apparently a honeybee mimic, and two are apparently 148 non-mimetic flies. Excluding the honeybee mimic (*Eristalis tenax*), we assigned the dipteran species 149 a rank according to the accuracy with which human volunteers perceived they mimicked P. 150 dominula. Twenty seven students at the University of Nottingham were shown one photograph 151 selected at random from three high-resolution colour photographs of *P. dominula*, and one of three 152 photographs (again, selected at random) of each of the six fly species. We used the best images of each species to which we had access, but the scale, lighting conditions etc. of these images varied as 153 154 was not always known. Volunteers were asked to rank the fly species in the photographs in order of 155 resemblance to P. dominula. We did not emphasise the importance of any particular cues/criteria in 156 scoring resemblance.

#### 157 Experiment 1: Prey Preference

158 A field experiment was carried out to determine if crab spiders distinguish among prey types, and to 159 establish the palatability of the defended model species (*P. dominula* and *A. mellifera*). Between 160 0900 and 1800 on 19 sunny days between April 3rd and May 5th 2008, live model and mimic species 161 were offered to *S. globosum* found foraging on flowers in the field and their subsequent response 162 recorded. Individuals of seven prey species (Table 1) were caught with a sweep net no more than 24 163 hours before the experiment and temporarily stored in specimen tubes. Immediately before the 164 experiment, each individual was stunned using carbon dioxide gas, its wings were removed, and a 165 piece of very fine metal wire (350 mm long and 0.125 mm in diameter) tied between its head and thorax. The removal of the wings ensured that the prey were easier to manipulate, and minimised 166 167 the extent of any auditory or behavioural cues which might influence spider behaviour. Each 168 individual was suspended on its wire from a 30 cm wooden stick and offered to a different spider (n 169 = 180; 139 females and 41 males). An individual of a randomly selected prey taxon was "dangled" 5 -170 10 mm above the centre of a haphazardly selected flower with a resident spider. Random selection 171 of prey types meant that sample sizes were not necessarily equal. Individuals were dangled for 5 min 172 or until they were killed by the spider. In addition to whether prey were killed and consumed by 173 spiders, seven distinct spider behaviours were recorded during trials (Table 2). During the course of 174 Experiment 1, it became clear that "retreat" actually encompassed two distinct behaviours, one of 175 which was subsequently redefined "bungee".

176 Experiment 2: Can Spiders Learn Aversion to Mimetic Prey?

In order to control the previous experiences and hunger of the crab spiders, all subsequent
experiments were carried out under laboratory conditions at our field station. Spiders used in
laboratory experiments were collected from flowers in the field, stored individually in 9 cm Petri
dishes and kept on a natural light:dark cycle. Prior to their use in experiments, they were starved for
at least four days to increase their motivation to hunt.

182 The first laboratory experiment tested whether prior experience with an aposematic, aversive model 183 species affected spider behaviour when subsequently presented with mimetic and non-mimetic 184 palatable prey. Field observations and the results of Experiment 1 indicated that P. dominula and 185 hoverfly mimics of wasps were the most suitable species for this test: whilst S. globosum readily 186 attacks and consumes A. mellifera, it rarely if ever attacks P. dominula, and in fact often flees from a 187 flower upon encountering this species. Thus, we tested 1) whether spiders exposed to P. dominula 188 were less likely to attack wasp mimics subsequently, and 2) whether any such acquired aversion to 189 wasp mimics was stronger towards hoverfly species which are considered by humans to be more 190 accurate mimics.

Over 27 days in April – June 2008 and March – May 2009, spiders (n = 259; 188 females and 71 191 192 males) were randomly allocated to either "wasp" or "no-wasp" treatments and placed in a clean 193 Petri dish under a desk lamp with a 60W bulb for 10 min. Individuals in the wasp treatment were 194 housed with a live adult P. dominula (with wings removed). In a few cases, spiders were injured or 195 killed by wasps, in which case they were discarded. After 10 min, spiders were removed from dishes 196 and placed onto a fresh flower of Chrysanthemum segetum standing in a sample tube filled with 197 water, in a Perspex experimental arena (30 cm x 20 cm x 20 cm). C. segetum was chosen because 198 crab spiders were often seen foraging on it the field and it did not wilt easily. The spider was given 199 10 min to settle on the flower, after which a fly specimen from one (randomly selected for each trial) 200 of six species (four wasp mimics and two non-mimics; see Table 1) was dangled above the centre of 201 the flower as above. Random selection of treatments for individual spiders meant that sample sizes 202 were not necessarily equal. Spider behaviours were recorded as before (Table 2). On rare occasions, 203 spiders did not settle on the flower; these individuals were discarded. For this experiment, prey 204 were frozen until dead immediately prior to being presented to spiders, in order to remove 205 mechanosensory stimuli. Preliminary tests revealed that spiders were equally likely to attack live and 206 dead prey.

207 Experiment 3: Visual Cues

208 The next experiment was designed to establish whether S. globosum relies on visual cues whilst 209 hunting and whether, after experience with a wasp, they show aversion towards visual signals 210 designed to mimic the black and yellow banding on vespid abdomens. Over 15 days in May 2008 and 211 in March and April 2009, we presented spiders with dead flies and artificial "prey" consisting of 10 212 mm long cylindrical pieces of modelling clay (Staedtler<sup>®</sup> Fimo<sup>®</sup>). Both real and artificial prey were 213 manipulated to produce mimetic and non-mimetic forms. Artificial prey were either black clay with 214 black stripes, yellow clay with black stripes or yellow clay with yellow stripes. All stripes were 215 painted using a fine brush and Revell® enamel model paint (yellow matt #15 and black matt #8). A groove cut approximately 2 mm from one end of each cylinder acted as the 'neck' to allow metal 216 217 wire to be fastened in place (as above). The flies used were the hoverfly C. intermedium, which was 218 deemed the most accurate mimic of P. dominula by human volunteers, and the non-mimetic fly F. 219 canicularis (Table 1). Their wings and legs were removed. Two "altered" forms of C. intermedium 220 were also used for comparison with artificial prey, and to control for possible effects of paint on 221 spider behaviour: they had either their yellow markings or their black markings painted over using 222 black paint (as above). Spiders (n = 98; 69 females and 29 males) were exposed to wasps as in the 223 wasp treatment in Experiment 2, and then transferred to flowers in the experimental arena. They 224 were offered artificial prey of a randomly selected type, as above, for 3 min or until the spider 225 attacked. Spider behaviour was recorded as before (Table 2). Random selection of the prey type for 226 each individual spider meant that sample sizes were not necessarily equal.

227 Experiment 4: Olfactory Cues

The final experiment was designed to determine whether *S. globosum* uses olfactory cues whilst hunting, and whether there is olfactory mimicry in the apparently accurate visual wasp mimic *C. intermedium*. All prey in this experiment were cylindrical pieces of black modelling clay. Each cylinder was randomly assigned an odour treatment: wasp, honeybee, wasp mimic or nothing. 232 Odour was transferred using a similar method to that used by Wood and Ratnieks (2004). Each 233 cylinder was placed in a sample tube with a live wasp (P. dominula), honeybee (A. mellifera), wasp 234 mimic (C. intermedium) or nothing, and shaken for ten seconds. We shook the tube sufficiently to 235 prevent the live animals from avoiding contact with the artificial prey, but not so vigorously that they 236 were obviously harmed. The experiment was carried out over 15 days in April 2010. Each spider (n = 237 238; 161 females and 77 males) was randomly assigned to a "wasp" or "no-wasp" treatment, and 238 exposed to P. dominula for 10 minutes as in Experiment 2. Spiders were then transferred to a flower 239 in the experimental arena and offered a prey cylinder from a randomly chosen odour treatment for 240 three minutes, or until it was attacked. Behaviours exhibited by the spiders during trials were 241 recorded as before (Table 2).

242 Statistical Analysis

243 For analysis, we characterised spider responses to prey in two different ways. First, where possible, 244 we considered whether or not prey were killed in a trial as a binary response variable. In 245 experiments with dead or artificial prey, we considered whether or not prey were attacked at least 246 once, instead of whether they were killed. Second, we considered the frequency with which each 247 different behaviour (Table 2) occurred in each trial. Because the frequencies of some behaviours 248 were clearly correlated, we used Principal Components Analysis (PCA) to derive a smaller number of 249 uncorrelated response variables with which to describe these data. We extracted all Principal 250 Components (PCs) with Eigenvectors of greater than 1.0 for use in subsequent analysis.

The effects of spider sex, prey taxon or type, treatment (exposure to *P. dominula*) and prey mimetic accuracy on the two types of response variable were analysed using either binomial or Gaussian generalised linear models (GLMs) or linear mixed effects models (GLMMs). For all experiments, we fitted GLMs, in which spider sex and prey taxon/type were fixed factors, and there were no random effects. Where appropriate, treatment was also fitted as a fixed factor. In Experiments 1 and 2, we explicitly tested the effect of mimetic accuracy using a GLMM, with prey taxon as a random factor

257	and prey mimetic accuracy (ranked) as a fixed covariate. Model fitting and simplification followed
258	Zuur et al. (2009). The significance of fixed effects was assessed by testing the effect of deleting
259	terms sequentially, starting with the highest order interactions, from a saturated model with
260	Likelihood Ratio, F-ratio or Chi-squared tests. PCA and model fitting was done using the packages
261	glmer and stats in R Version 3.0.1 (R Core Team, 2015). Post hoc comparisons of prey taxa were
262	performed using the package phia. Where relevant, planned comparisons were applied with a
263	custom contrast matrix.

#### Results

### 266 Experiment 1: Prey Preference

267 When offered prey in the field, spiders were significantly more likely to kill and consume some taxa than others (binomial GLM:  $\chi^2_7$  = 89.434, p < 0.001; Figure 2). Spiders never killed *P. dominula*, but 268 269 readily consumed most dipteran prey, except for the honeybee mimic *E. tenax*. Female spiders were significantly more likely to kill prey (62.6 % of 139) than males (25 % of 40) ( $\chi^2_1$  = 8.6142, p = 0.003), 270 but there was no interaction between sex and prey taxon ( $\chi^2_7$  = 7.953, p = 0.337). Considering only 271 the data for the wasp mimics and control fly species, the effect of spider sex was no longer 272 significant (GLMM:  $\chi^2_1$  = 1.579, p = 0.209), spiders were not significantly more likely to kill less 273 accurate mimics ( $\chi^2_1$  = 0.295, p = 0.587), and there was no interaction between sex and accuracy ( $\chi^2_1$ 274 275 = 1.480, p = 0.224). 276 Principal Components Analysis (PCA) of spider behaviour yielded two Principal Components (PCs)

with Eigenvalues greater than one (Table 3). The first (PC1) correlated positively with the frequencies of all behaviours except "attack" and "left flower", and hence can be interpreted as a general index of activity exhibited by spiders when they encountered prey. The second (PC2) correlated positively with the number of attacks observed, and negatively with the number of times the spider left the flower upon which it was sitting, and hence contrasts the most positive and the most negative reactions by spiders to prey, with positive scores indicating positive reactions.

Spiders behaved differently towards different prey taxa. Scores on PC1 and PC2 were significantly different among prey taxa (PC1:  $F_{(7, 171)} = 6.204$ , p < 0.001; PC2:  $F_{(7,170)} = 2.591$ , p = 0.015), with spiders being most active and least likely to react positively to *P. dominula*, and least active but most likely to respond positively to the non-mimic *Fannia canicularis* and the relatively inaccurate wasp mimic *Sphaerophoria scripta* (Figure 3). There was a significant difference between male and female spiders in PC2 scores ( $F_{(1,170)} = 10.396$ , p = 0.002), with males being less positive towards prey (mean 289 = -0.460; SEM = 0.196) than females (mean = 0.132; SEM = 0.075), but not in PC1 scores ( $F_{(1,170)}$  = 0.013, p = 0.910). There was no interaction between the effects of sex and taxon on either PC (PC1: 290  $F_{(7,163)} = 1.717$ , p = 0.108; PC2:  $F_{(7,163)} = 1.879$ , p = 0.076). Considering only the data for the wasp 291 mimics and control fly species, male spiders were both more active (GLMM for PC1:  $\chi^2_1$  = 8.433, p = 292 0.004) and less positive towards prey (PC2:  $\chi^2_1$  = 7.302, p = 0.007). However, there was no significant 293 effect of mimetic accuracy on either measure of behaviour (PC1:  $\chi^2_1$  = 0.136, p = 0.713; PC2:  $\chi^2_1$  = 294 0.001, p = 0.999) and no interaction between accuracy and sex (PC1:  $\chi^2_1$  = 0.304, p = 0.581; PC2:  $\chi^2_1$  = 295 296 3.124, p = 0.077).

297 Experiment 2: Can Spiders Learn Aversion to Mimetic Prey?

298 When offered hoverflies with varying levels of mimetic accuracy, and non-mimetic flies, in the 299 laboratory, spiders with prior experience of P. dominula were slightly less likely to kill prey (63.6 % of 300 129) than spiders without such experience (72.3 % of 130), but this effect was not significant (binomial GLM:  $\chi^2_1$  = 2.354, p = 0.125). The probability that a spider killed a prey item was 301 significantly affected by taxon ( $\chi^2_1$  = 36.714, p < 0.001; see Figure 4), with the relatively accurate 302 303 mimic C. intermedium being killed least often, and the non-mimetic fly F. canicularis being killed most often. Spider sex did not significantly affect the probability of prey being killed ( $\chi^2_1$  = 3.145, p = 304 305 0.076). None of the interactions among the main effects on the probability that spiders killed a prey 306 item was significant (p > 0.3 in all cases). It is worth noting that, whilst most prey taxa were attacked 307 at similar frequencies in the two treatments, the most accurate mimic taxon was about half as likely 308 to be attacked by spiders if they had previously been exposed to P. dominula. However, when the 309 data were reanalysed with a mixed model, there was no significant effect of mimetic accuracy on the probability of prey being killed (binomial GLMM:  $\chi^2_1$  = 2.674, p = 0.102) and no significant 310 311 interactions involving accuracy (p > 0.1 in all cases). Tests of the other main effects and interactions 312 in the mixed model yielded qualitatively identical results to the GLM (results not shown).

PCA of spider behaviours in Experiment 2 produced three PCs with Eigenvalues greater than one (see
Table 3). The first two PCs were similar to those produced in Experiment 1: the first correlated
positively with all behaviours except "attack", and the second contrasted the frequency of attacks
with the frequency with which spiders left flowers. The third PC correlated most strongly (positively)
with "bungee", a behaviour which was not distinguished from "retreat" in Experiment 1.

318 Analysis which modelled the effect of prey taxon as a fixed factor (GLMs) showed that there were no 319 significant effects of prior exposure to *P. dominula*, spider sex or prey taxon, or their interactions, on 320 PC1 (Table 4; Figure 5). There were, however, significant main effects of prey taxon and treatment 321 on PC2 (Figure 6). Spiders behaved most positively towards S. ribesii and most negatively towards C. intermedium. In addition, spiders were less likely to attack prey, and more likely to flee them, if they 322 323 had previously been exposed to wasps. There were no significant interactions involving these main 324 effects on PC2. The behaviours captured by PC3 showed a complex response to the different 325 explanatory variables, with a significant three-way interaction in addition to a main effect of taxon 326 (Fig S1, supplementary information). In general, spiders were least likely to "bungee" (i.e. had the 327 lowest PC3 scores) in response to C. intermedium, and most likely to exhibit this behaviour in 328 response to S. ribesii. The significant three-way interaction reflects contrasting patterns in male and 329 female PC3 scores, particularly towards S. ribesii, S. scripta and E. balteatus: prior exposure to wasps 330 had a relatively positive effect on the male propensity to "bungee" in response to S. ribesii and E. 331 balteatus, and a relatively negative effect on the male propensity to "bungee" in response to S. 332 scripta, when compared to the equivalent effects of exposure on females.

The mixed model analysis, which included a fixed effect of mimetic accuracy, did not reveal any significant main effects on any of the principal components, but did indicate a significant interaction between treatment and mimetic accuracy: spiders which had been previously been exposed to the aversive model species were relatively more active in response to more accurate mimics, in comparison with less accurate or non-mimetic prey (Figure 5). When spiders previously exposed to *P. dominula* were offered real and artificial prey whose visual appearance was more or less similar to the aversive model species, the probability that they would attack was significantly affected by prey type (binomial GLM:  $\chi^2_6 = 21.173$ , p = 0.002; Figure 6). Unmanipulated hoverfly mimics with black and yellow stripes (*C. intermedium*) and stripy artificial prey were the least likely to be attacked. According to planned comparisons, artificial prey without stripes were significantly more likely to be attacked than those with stripes. Hoverflies with their

345 yellow or black stripes painted black were more likely to be attacked than unpainted equivalents,

346 although not significantly so. Unmanipulated, non-mimetic control flies were the prey type most

347 likely to be attacked. Spider sex (GLM:  $\chi^2_1 = 0.034$ , p = 0.854), and the interaction between sex and

348 prey type (GLM:  $\chi_6^2$  = 8.212, p = 0.223), did not significantly affect the probability of attack.

349 As in Experiment 2, PCA produced three PCs with Eigenvalues greater than one, the first two of 350 which had similar correlations with the raw variables describing spider behaviour (Table 3). None of these PCs was significantly affected by prey type (Gaussian GLM for PC1: F<sub>6,89</sub> = 0.941, p = 0.470; PC2: 351 352  $F_{6,89}$  = 1.967, p = 0.079; PC3:  $F_{6,89}$  = 0.750, p = 0.611), although PC2 (contrasting the propensity to 353 attack with the tendency to leave the flower) showed patterns consistent with the above analysis of 354 spider attacks: stripy artificial prey scored lower than those without stripes, and the same was true 355 for hoverflies, whilst unmanipulated control flies had the most positive scores (Fig S2, 356 supplementary infromation). There were no significant effects of spider sex (F<sub>1,88</sub> < 3.0, p > 0.05) or

the interaction between sex and prey type ( $F_{6,82} < 1.5$ , p > 0.1) on PC scores.

358 Experiment 4: Olfactory Cues

When offered artificial prey bearing the odour of models (*P. dominula* and *A. mellifera*) and mimics (*C. intermedium*), or no odour, in the laboratory, there were no significant effects of prior experience with *P. dominula* (binomial GLM:  $\chi_{1}^{2} = 1.689$ , p = 0.194), odour type ( $\chi_{3}^{2} = 1.609$ , p = 0.657) or spider sex ( $\chi^2_1$  = 1.568, p = 0.211) on the probability that spiders would attack. Neither were there any significant effects of any interactions among these effects on the probability that spiders would attack prey ( $\chi^2_{1 \text{ or } 3} < 5.0$ , p > 0.1).

As in Experiments 2 and 3, PCA produced three PCs with Eigenvalues greater than one, the first two of which had similar correlations with the raw variables describing spider behaviour (Table 3). None of these PCs was significantly affected by odour type or prior experience with *P. dominula* (Table 5). However, PC1, but not PC2 and PC3, was significantly affected by spider sex, with male spiders scoring more highly, indicating higher frequencies of most behaviours (especially display, retreat and approach) than were seen in females. There were no significant interactions among the effects of prior experience, odour type and sex on any of the PCs.

#### 372 Discussion

373 The results of our experiments demonstrate that the crab spider *S. globosum* has clear prey

374 preferences, and suggest that those preferences are influenced by the visual appearance of prey, but

not by olfactory cues. In addition, there was limited evidence that prior experience with an aversive

376 model species altered spider behaviour towards more accurate Batesian mimics of that model.

377 Overall our results provide support the idea that invertebrate predators can play a role in the

evolution of the visual appearance of their prey, and shed some light on the hypothesis that

379 differences between invertebrate and vertebrate sensory and cognitive processes result in relaxed

380 selection on mimetic perfection in prey which are subject to predation by invertebrates.

381 Prey preferences

The existence and nature of prey preferences in crab spiders are poorly reported. The large, flowerdwelling species *Misumena vatia* apparently shows very little discrimination among prey types, and the frequencies of prey taxa in its diet are thought simply to reflect variation in prey availability (Morse, 2007). By contrast, our results show that *S. globosum* readily distinguished among prey 386 species in the field, completely avoiding *P. dominula*, whilst killing and consuming nearly all 387 individuals of some dipteran taxa. S. globosum therefore has the potential to exert selection 388 pressure on prey phenotypes via the cues it uses to identify prey types. Other studies have shown 389 that even when spiders are able to discriminate among prey types, this discrimination may not affect 390 selection, because even rejected prey were left irreparably injured or dead (see Vasconcellosneto 391 and Lewinsohn, 1984). In the case of S. globosum, however, our observations showed that none of 392 the prey which were bitten were later rejected, and all rejected prey were left unharmed. Therefore, 393 especially given its abundance and likely impact on prey populations (Ibarra and Reader, 2014; 394 Reader et al., 2006), the results of our experiments suggest that S. globosum does indeed have the 395 potential to select for visual or other traits in aposematic and mimetic prey.

396 The putative model wasp species *P. dominula* was never attacked in the field experiment.

397 Furthermore, encountering *P. dominula* caused the spiders to become very active, often leaving the 398 flower completely during the prey presentation period. This suggests that P. dominula was aversive 399 to S. globosum, either because it is unprofitable, distasteful, or dangerous. Birds may avoid vespid 400 wasps because their abdomens are distasteful (Mostler, 1935), but circumstantial evidence points 401 away from this explanation for avoidance of P. dominula by S. globosum: while other spiders have 402 been shown to attack unpalatable prey before rejecting it moments later (Vasconcellosneto and 403 Lewinsohn, 1984), we have only seen S. globosum attack P. dominula once (during exposure in the 404 laboratory in Experiment 2), and on this occasion the spider appeared to feed unperturbed until the 405 wasp was completely consumed. Instead, the main deterrent associated with P. dominula appears to 406 be its aggressive and/or predatory nature. In the laboratory, it frequently bit or attempted to sting S. 407 *globosum*, killing spiders on several occasions, and in the field *P. dominula* is often seen hunting 408 invertebrate prey on or around flowers which harbour S. globosum. It is therefore high plausible that 409 learned or evolved avoidance of *P. dominula* is adaptive in *S. globosum*.

410 In contrast to the results for P. dominula, the other aposematic, defended insect we offered to S. 411 *globosum* in Experiment 1 – the honeybee – was readily attacked and consumed, a fact which helps 412 to explain the levels of mortality that spiders impose on honeybees in our field site (Reader et al., 413 2006). For this reason, we excluded the honeybee and its mimic *E. tenax* from subsequent 414 experiments in which we were interested in responses to mimicry of aversive prey. The honeybee's 415 defences are apparently not a threat to S. globosum, which underlines the likely importance of P. 416 dominula's aggressive predatory behaviour, rather than the possession of a sting per se, in 417 determining spider prey choice. The almost complete avoidance of *E. tenax*, which to humans is a 418 good mimic of the honeybee, in Experiment 1 is puzzling. It suggests that, although visual 419 appearance may play a role in some circumstances (as in Experiment 3, for example), other factors 420 affect spider prey choice significantly. Which cues were used to identify *E. tenax* as unpalatable, and 421 the reason why it is unpalatable, remain a mystery which only further experimentation can unlock.

422 In two of our four experiments, sex had a significant effect on spider responses to prey. There was 423 no evidence that prey preferences were different between males and females, but males were in 424 general less likely to attack prey in the field, and more active (rather "skittish") when presented with 425 artificial prey in the laboratory. These differences could be explained by the pronounced sexual size 426 dimorphism (Blanckenhorn, 2005) that is seen in many spiders, including S. globosum (full grown 427 females are about seven times the mass of males; Corcobado et al., 2010). Typically, smaller male 428 spiders have a different diet from larger females (Walker and Rypstra, 2002). Furthermore, males 429 often spend the majority of the time searching for and guarding females and invest less energy and 430 time in foraging for themselves (e.g. Givens, 1978) than females, whose reproductive fitness is 431 closely related to foraging success (Walker and Rypstra, 2002). As a result, male S. globosum may 432 have been more interested in fleeing the confines of the experimental arena than they were in 433 finding a meal. Their smaller body size did not, however, obviously constrain prey choice: just like 434 females, males showed a relative preference for the largest popular prey type (honeybees) 435 compared with some of the smaller prey in Experiment 1, for example.

#### 436 Responses to Batesian mimicry

437 Our results provide some support for the hypothesis that *S. globosum* perceives and responds to 438 Batesian mimicry in its prey. Results from the laboratory (Experiment 2) showed that, overall, spider 439 attack rates and other measures of behaviour towards hoverflies which mimic wasps did not 440 correlate strongly with mimetic accuracy, as perceived by humans. However, recent prior exposure 441 to the aversive model P. dominula resulted in some changes in behaviour which were most marked 442 towards the hoverflies which most closely resemble the model. Whilst the relatively large (c. 50 %) 443 decline in the frequency of attacks by spiders exposed to the model on the best mimic (C. 444 *intermedium*) was not significantly different from the change in the attack rate on other prey 445 species, there was a significant interaction between the effects of mimetic accuracy and exposure to 446 the model on spider activity levels. The high levels of spider activity which characterised their 447 response to the model species in Experiment 1 were mirrored in exposed spiders offered relatively 448 accurate mimics (especially C. intermedium) in Experiment 2. One explanation for this pattern is that 449 exposed spiders generalised from learned responses to wasp cues, resulting in them avoiding or 450 trying to escape from those particular prey taxa which humans perceive to be accurate Batesian 451 mimics. Hence, although the statistical support for the observed pattern is somewhat equivocal, and 452 we do not know what the effects on prey fitness would be in the wild, our data do provide some 453 tentative suggestions that spiders might respond to Batesian mimicry (see also Tyshchenko, 1961). 454 There are, however, alternative explanations for the observed patterns. Rather than facilitating 455 learning about wasp-related cues, a recent encounter with a potential predator may "prime" S. 456 globosum for danger, making it more likely to exhibit innate avoidance behaviours in subsequent 457 encounters with certain prey types. This priming might not be specific to *P. dominula*, and might 458 have nothing to do with aposematism or mimicry. We would have to repeat the experiment with a control treatment in which spiders were exposed to a different insect (e.g. one which was not 459 460 aposematic, and not dangerous) in order to explore the possibility further.

461 Although we found some evidence that mimetic accuracy as perceived by humans may affect spider 462 behaviour, there are several reasons why we might expect spiders to rank mimics differently from 463 vertebrates. In addition to differences in sensory and cognitive capabilities, spiders probably also 464 view their insect prey from a different angle from that typically experienced by birds or humans. In 465 our study, humans evaluated the accuracy of mimics from photographs taken from above, whilst 466 spiders were beneath their prey as they were "dangled" onto flowers. Since prey appear very 467 different from below, this could substantially change the accuracy of any perceived mimicry from 468 the spider's perspective, but the significance of any such change remains unknown without testing 469 the effect of prey orientation on human and spider behaviour.

470 There were other limitations to the ranking exercise we used to assess mimetic accuracy of the 471 insect taxa we used in our experiments. For example, we were not able to standardise the scale of 472 the images we showed to human volunteers, and hence reliable information about body size was 473 not available to them, in contrast to the situation for spiders in the experiments. Overall, therefore, 474 we are cautious about the interpretation of our analyses which included mimetic accuracy as a 475 predictor, and we think the most important message from our study is that spiders responded 476 differently to different prey taxa and different artificial prey types in a way which highlights the 477 potential of spiders as agents of selection on colour patterns in mimetic and aposematic insects.

478 The importance of visual, olfactory and other cues

The results show that *S. globosum* relies at least partly on vision when hunting. While the preferences demonstrated in the field (Experiment 1) may have been influenced by olfactory and/or mechanosensory information associated with live prey, Experiment 3 showed that spiders also discriminated among artificial prey that varied only in colour. The results from Experiment 3 also suggest that *S. globosum* responds to typical aposematic signals: following previous wasp experience, the spiders attacked black and yellow striped artificial prey significantly less often than the completely yellow and completely black artificial prey. It remains unclear whether the important cue here was variation in hue (yellow versus black) or simply luminance (contrast between dark and
light stripes), but evidence for true colour vision in crab spiders (Thomisidae) is limited (Insausti et
al., 2012).

There was no evidence that *S. globosum* distinguished among the different odour treatments in Experiment 4. The results thus suggest that crab spiders do not utilise olfactory signals to detect the aversive model *P. dominula* or its mimcs. This could be because the olfactory signals of wasps have not evolved under selection by predators, and are not as readily learned and remembered as aposematic colour signals are thought to be (Stevens, 2007; Svadova et al., 2009). Alternatively, *S. globosum* may not rely heavily on olfactory signals while foraging, as appears to be the case in the closely related crab spider, *Misumena vatia* (Morse, 2007).

496 The laboratory experiments presented here were designed to eliminate behavioural, auditory and 497 mechanosensory stimuli emitted by prey. Under natural conditions, these stimuli may contribute to 498 decision making by crab spiders, and hence they may play a role in determining the adaptive value of 499 wasp mimicry by hoverflies. There is evidence of behavioural mimicry in some hoverfly species 500 (Rotheray and Gilbert, 2011), but the only published evidence suggests that hoverflies do not mimic 501 the sounds of their hymenopteran models (Rashed et al., 2009). Observations of the behaviour of 502 the *M. vatia* suggest that it depends largely on mechanoreceptors when hunting (Morse, 2007), but 503 we know little about variance in the kind of vibrations produced among the potential prey of crab 504 spiders. In order to understand fully the potential of predators such as crab spiders to influence the 505 evolution of prey morphology and behaviour, further investigation of the importance of cues in each 506 sensory modality is required.

507 Conclusion

Overall, this study provides evidence that spiders have the ability to differentiate among prey using
visual cues, and therefore that they have the potential to exert selection on prey morphology,

510 including perhaps aposematic and mimetic colour patterns. Previous studies of Batesian mimicry in 511 insects have tended to assume that the predators responsible for the evolution of warning patterns 512 and their mimetic equivalents are vertebrates. Given that invertebrates are important predators of 513 flower-visiting insects and several other groups containing aposematic and mimetic species, our 514 results suggest that the evolutionary significance of selection on aposematic and mimetic patterns 515 by invertebrate predators is worthy of more detailed consideration. The very different sensory and 516 cognitive abilities of invertebrate predators such as spiders could easily result in patterns of 517 selection which are not well predicted by human or avian behaviour, and hence our results are 518 broadly consistent with the hypothesis that the perceived degree of perfection of a Batesian mimic 519 may be in "the eye of the beholder" (Cuthill and Bennett, 1993). Crucially, in our community of 520 models and mimics, and in most others, the relative contributions of different types of predator to 521 selection on mimetic patterns remains unknown. The phenotype of a Batesian mimic should 522 represent a net evolved response to selection imposed by all predators. Careful consideration of the 523 importance and selectivity of predation from all sources is therefore needed to understand properly 524 how and why mimetic signals appear as they do.

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Figure 1. Nine insect species offered to crab spiders in experiments. A) The honeybee (*Apis mellifera*)
and B) its mimic (*Eristalis tenax*). C) The paper wasp (*Polistes dominula*). Wasp mimics in descending
order of accuracy, as perceived by humans: D) *Chrysotoxum intermedium*, E) *Syrphus ribesii*, F) *Sphaerophoria scripta*, and G) *Episyrphus balteatus*. Non-mimetic flies: H) *Sarcophaga carnaria* and I)

Fannia canicularis. Images A, B and E are courtesy of Chris Taylor. Other images were taken by RM.

Figure 2. Differences in the proportion (+/- 95% Cl) of prey killed and consumed by spiders offered
different prey taxa in the field. Letters above bars indicate results of post hoc tests: species which
share a latter were not significantly different (p > 0.05).

641 Figure 3. Differences in behaviour of spiders towards different prey taxa offered to them in the field 642 in Experiment 1. Behavioural responses are characterised by means (+/- SEM) of two Principal 643 Components (PC1 and PC2) derived from Principal Components Analysis of the frequencies of seven 644 different behaviours observed in the field (see Table 3). High scores on PC1 indicate high frequencies of most behaviours except "attack" and "left flower". High scores on PC2 indicate high frequencies 645 646 of "attack" and low frequencies of "left flower". Post hoc tests for PC1 showed that P. dominula and 647 *E. tenax* were significantly different from all other taxa (p < 0.05), but that there were no pairwise 648 differences amongst other taxa. Post hoc tests for PC2 showed that *P. dominula* was significantly 649 different from all other taxa, but that there were no pairwise differences amongst other taxa.

Figure 4. The effects of previous exposure to the aversive model *P. dominula*, and prey mimetic
accuracy, on the probability (+/- 95% CI) that spiders attacked prey of various taxa offered to them in
the laboratory in Experiment 2.

Figure 5. The effects of previous exposure to the aversive model *P. dominula*, and prey mimetic
accuracy, on spider behaviour when offered different prey taxa in the laboratory in Experiment 2.
Spider behaviour is described by mean PC1 scores (+/- SEM) derived from Principal Components

Analysis of the frequencies of eight different behaviours observed in the field (see Table 3), with high
scores indicating spiders which were more active and less likely to attack prey.

658 Figure 6. The effects of previous exposure to the aversive model *P. dominula*, and prey taxon, on

659 spider behaviour when offered different prey taxa in the laboratory in Experiment 2. Spider

behaviour is described by mean PC2 scores (+/- SEM) derived from Principal Components Analysis of

the frequencies of eight different behaviours observed in the field (see Table 3), with high scores

indicating spiders which were more likely to attack prey, and less likely to flee from them.

Figure 7. The effects of prey type on the proportion of spiders (+/- 95% CI) that attacked real and

artificial prey in the laboratory in Experiment 3. Prey types were: Natural Mimic (the accurate wasp-

665 mimic C. intermedium; n = 13), Black Mimic (C. intermedium with its yellow stripes painted black; n =

11), Control Mimic (*C. intermedium* with its black stripes painted black; n = 9), Stripy Artificial (yellow

667 clay with black stripes; n = 16), Yellow Artificial (yellow clay with yellow stripes; n = 16), Black

668 Artificial (black clay with black stripes; n = 18) and Non-mimic (the non-mimetic fly *F. canicularis*; n =

15). Planned comparison tests are indicated above the bars for pairs of prey types of particular

670 interest (NS: p > 0.05; \* p < 0.05).

- Table 1. Prey types used, their mimetic status, the experiments in which they were used, and their
- 672 ranked mimetic accuracy as perceived by humans (see main text).

Species / prey type	Mimetic status	Experiments	Mimetic Rank (1 = most
			accurate)
Polistes dominula (paper wasp) (L. 1758)	Model	All	NA
Chrysotoxum intermedium (Meigen	Wasp mimic	All	1
1822)			
Syrphus ribesii (L. 1758)	Wasp mimic	1&2	2
Sphaerophoria scripta (L. 1758)	Wasp mimic	1 & 2	3
Episyrphus balteatus (De Geer 1776)	Wasp mimic	2	4
Apis mellifera (honeybee) (L. 1758)	Model	1	NA
Eristalis tenax (L. 1758)	Honeybee	1	NA
	mimic		
Sarcophaga carnaria (L. 1758)	Not mimetic	1&2	5
Fannia canicularis (L. 1758)	Not mimetic	1, 2 & 3	6
C. intermedium paint control	Wasp mimic	3	NA
C. intermedium black	Not mimetic	3	NA
Artificial black	Not mimetic	3 & 4	NA
Artificial yellow	Not mimetic	3	NA
Artificial back and yellow stripes	Wasp mimic	3	NA

- Table 2. Behaviours performed by *S. globosum* during experiments. Note that retreat and bungee
- 675 were not differentiated in Experiment 1.

Behaviour	Description
Leave Flower	Leaving the plant completely and not returning
Bungee	Jumping from the flower and dangling on a line of silk
Retreat	Moving away from the prey (and, in Experiment 1 only, bungeeing)
Display	Spreading and lifting fore-limbs while orientated towards prey
Approach	Orientation and movement towards the prey
Tickle	Gently touching the prey with its front legs
Grapple	Frantic touching of the prey and occasional jumping
Attack	Mounting and biting of the prey

Table 3. Results of Principal Components Analysis of behavioural responses by spiders to prey offered in the field in Experiment 1, and in the laboratory in

678 Experiments 2, 3 and 4. Correlation coefficients are shown indicating the strength and direction of the relationships between extracted Principal

679 Components (with Eigenvectors > 1.0) and the original variables describing the frequencies with which particular behaviours (see Table 2) were observed.

680 Note that retreat and bungee were not differentiated in Experiment 1.

	Experime	ent 1	Experime	ent 2		Experime	ent 3		Experime	ent 4	
Original variable	PC1	PC2	PC1	PC2	PC3	PC1	PC2	PC3	PC1	PC2	PC3
Display	0.751	0.095	0.551	0.347	-0.044	0.772	0.268	0.230	0.821	0.304	0.097
Retreat	0.884	0.003	0.894	0.025	-0.114	0.849	-0.078	0.019	0.836	-0.270	0.053
Approach	0.874	0.145	0.620	0.190	0.550	0.749	-0.139	-0.01	0.836	-0.136	-0.070
Tickle	0.704	-0.189	0.717	0.140	-0.123	0.471	-0.160	-0.636	0.436	0.106	-0.174
Grapple	0.409	0.261	0.511	0.167	-0.498	0.634	0.402	0.297	0.698	0.395	0.219
Attack	-0.196	0.729	-0.459	0.713	0.059	-0.194	0.661	0.405	-0.109	0.809	-0.141
Bungee	NA	NA	0.236	-0.235	0.789	0.198	-0.751	0.285	0.332	-0.441	-0.452
Left flower	0.041	-0.745	0.165	-0.812	-0.193	-0.097	-0.621	0.541	-0.010	-0.213	0.839
Variance explained	40.17 %	17.44 %	31.96 %	17.85 %	15.54 %	32.34 %	20.87 %	13.55 %	35.88 %	15.57 %	12.79 %

Table 4. Results of analysis of the effects of spider sex, prey taxon or mimetic accuracy, and treatment (whether spiders were exposed to a wasp before encountering prey), and their interactions, on Principal Components (PCs) describing the behavioural responses of spiders to prey in the laboratory in Experiment 2 (see Table 3). Models were fitted with a Gaussian error structure, and included either a fixed effect of prey taxon (GLMs), or a fixed effect of mimetic accuracy and a random effect of prey taxon (GLMMs). Effects were assessed by F tests or Likelihood Ratio (LR) deletion tests during model

686 simplification. Significant p values are highlighted in bold.

	GLM with fixed effect of taxon G								GLMM with fixed effect of accuracy									
	PC1			PC2			PC3			PC1				PC2			PC3	
Term	F	df	р	F	df	р	F	df	р	LR	df	р	LR	df	р	LR	df	р
Sex	0.061	1, 252	0.806	0.664	1, 251	0.416	3.782	1, 252	0.053	1.036	1	0.309	1.037	1	0.309	0.755	1	0.385
Taxon/Accuracy	0.924	5, 253	0.466	3.562	5, 253	0.004	5.573	5, 253	< 0.001	1.639	1	0.201	0.727	1	0.394	2.084	1	0.149
Treatment	0.062	1, 251	0.803	5.055	1, 252	0.025	0.498	1, 251	0.481	3.410	1	0.065	1.489	1	0.222	2.369	1	0.124
Sex x Taxon/Accuracy	0838	5, 240	0.524	0.466	5, 241	0.802	1.089	5, 246	0.367	0.756	1	0.385	2.301	1	0.129	0.327	1	0.568
Sex x Treatment	0.138	1, 245	0.710	0.001	1, 240	0.995	1.820	1, 241	0.110	0.162	1	0.687	0.027	1	0.869	2.394	1	0.122
Taxon/Accuracy x Treatment	1.213	5, 246	0.304	0.649	5, 246	0.663	0.798	5, 240	0.373	5.131	1	0.024	0.149	1	0.699	2.686	1	0.101
Sex x Taxon/Accuracy x Treatment	2.235	5, 235	0.052	0.910	5, 235	0.968	4.520	5, 235	< 0.001	2.348	1	0.125	0.206	1	0.650	0.858	1	0.354

Table 5. Results of analysis of the effects of spider sex, prey odour and treatment (whether spiders
were exposed to a wasp before encountering prey), and their interactions, on Principal Components
(PCs) describing the behavioural responses of spiders to prey in the laboratory in Experiment 4 (see
Table 3). Effects were assessed by F-tests at deletion during simplification of a Gaussian general

- 692 linear model.

	PC1			PC2			PC3			
Term	F	df	р	F	df	р	F	df	р	
Sex	13.745	1,236	< 0.001	0.245	1,236	0.622	1.638	1,236	0.202	
Odour	1.320	3,232	0.269	0.031	3,232	0.993	0.855	3,233	0.465	
Treatment	2.379	1,235	0.124	1.023	1,235	0.313	0.416	1,229	0.520	
Sex x Odour	1.979	3,226	0.118	0.998	3,226	0.395	2.022	3,230	0.116	
Sex x Treatment	0.014	1,225	0.905	0.105	1,225	0.746	3.223	1,228	0.074	
Odour x Treatment	1.328	3,229	0.266	0.961	3,229	0.412	0.432	3,225	0.730	
Sex x Odour x Treatment	2.025	3,222	0.111	0.871	3,222	0.457	4.130	3,222	0.243	

















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707 Supplementary information

#### 708 Figure legends

- Figure S1. The effects of previous exposure to the aversive model *P. dominula*, and prey taxon, on the behaviour of male (a) and female (b) spiders, when offered
- 710 different prey taxa in the laboratory in Experiment 2. Spider behaviour is described by mean PC3 scores (+/- SEM) derived from Principal Components Analysis of
- the frequencies of eight different behaviours observed in the field (see Table 3), with high scores indicating spiders which were more likely to "bungee" from the
- 712 flower on which they were foraging.
- 713 Figure S2. The effects of prey type on the behaviour of spiders towards real and artificial prey in the laboratory in Experiment 3. Prey types were: Natural Mimic
- 714 (the accurate wasp-mimic *C. intermedium*; n = 13), Black Mimic (*C. intermedium* with its yellow stripes painted black; n = 11), Control Mimic (*C. intermedium* with
- its black stripes painted black; n = 9), Stripy Artificial (yellow clay with black stripes; n = 16), Yellow Artificial (yellow clay with yellow stripes; n = 16), Black Artificial
- 716 (black clay with black stripes; n = 18) and Non-mimic (the non-mimetic fly *F. canicularis*; n = 15). Spider behaviour is described by mean PC2 scores (+/- SEM)
- 717 derived from Principal Components Analysis of the frequencies of eight different behaviours observed in the field (see Table 3), with high scores indicating spiders
- which were more likely to more likely to attack prey, and less likely to flee from them.
- 719











Figure S2.