How predator hunting-modes affect prey behaviour: Capture 1 deterrence in *Drosophila melanogaster* 2 3 Abhijna Parigi<sup>1,2,3</sup>, Cody Porter<sup>2,4</sup>, Megan Cermak<sup>2</sup>, William R. Pitchers<sup>1,3</sup> and Ian Dworkin<sup>1,2,3\*</sup> 4 5 6 **Affiliations:** 7 1 Program in Ecology, Evolutionary Biology, and Behaviour, Michigan State University, East 8 Lansing, MI 48824, USA 9 2 Department of Zoology, Michigan State University, East Lansing, MI 48824, USA 10 3 BEACON Centre for the Study of Evolution in Action, Michigan State University, East 11 Lansing, MI 48824, USA 12 4 Current address: Department of Zoology and Physiology, University of Wyoming. Laramie, 13 WY, USA. 14 \* Correspondence to Ian Dworkin: idworkin@msu.edu 15 16 Links to videos (also available in supplement) 17 Abdominal Lifting http://dx.doi.org/10.6084/m9.figshare.1185638 18 Stopping Behaviour http://dx.doi.org/10.6084/m9.figshare.1185639 19 Retreat http://dx.doi.org/10.6084/m9.figshare.1185640

**ABSTRACT** 

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

Hunting mode, the distinct set of behavioural strategies that a predator employs while hunting, can be an important determinant of the prev organism's behavioural response. However, few studies have considered how a predator's hunting mode influences antipredatory behaviours of a prey species. Here we document the influence of active hunters (zebra jumping spiders, Salticus scenicus) and ambush predators (Chinese praying mantids, *Tenodera aridifolia sinensis*) on the capture deterrence anti-predatory behavioural repertoire of the model organism, *Drosophila melanogaster*. We hypothesized that D. melanogaster would reduce overall locomotory activity in the presence of ambush predators, and increase activity with active hunters. First we observed and described the behavioural repertoire of D. melanogaster in the presence of the predators. We documented three previously undescribed behaviours- abdominal lifting, stopping and retreat- which were performed at higher frequency by D. melanogaster in the presence of predators, and may aid in capture deterrence. Consistent with our predictions, we observed an increase in the overall activity of D. melanogaster in the presence of jumping spiders (active hunter). However, counter to our prediction, mantids (ambush hunter) had only a modest influence on activity. We also observed considerable intra and inter-individual variation in response to both predator types. Given these new insights into *Drosophila* behaviour, and with the genetic tools available, dissecting the molecular mechanisms of anti-predator behaviours may now be feasible in this system.

## **INTRODUCTION**

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

Predation, a ubiquitous selective force, gives rise to and determines the nature of defensive traits in prey populations (Edmunds, 1974; Juliano & Gravel 2002; Goslin & Rodd, 2007; Langerhans, 2007; Lima & Dill, 1990; Sansom, Lind & Cresswell, 2009). Predator hunting-modes, i.e., the set of behaviours that predators employ to pursue and capture their prey (Schoener, 1971; Huey & Pianka 1981; Preisser, Orrock & Schmitz 2007), have been shown to induce distinct prey responses (Schmitz, 2008) that in turn influence the productivity of ecological communities. In habitats dominated by active hunters there is lower species evenness and higher above-ground net primary productivity compared to habitats dominated by ambush hunters (Schmitz, 2008). The authors suggest the observed differences in prey productivity to be driven by hunting mode specific tradeoffs between foraging and seeking refuge. Although studies often describe the effects of predators on prey traits (i.e. DeWitt, Robinson & Wilson, 2000; Reznick, Butler & Rodd, 2001; Relyea, 2001), it is rare for the role of predator hunting-mode to be explicitly considered. Here we investigate segregating differences in the anti-predatory behavioural repertoire of the fruit fly, *Drosophila melanogaster*, in response to two predator species differing in hunting modes. Based on (Schmitz, 2008), we predicted that fruit flies, in the presence of a familiar predator, would exhibit hunting-mode specific modifications in activity levels. We used D. melanogaster because, although it is one of the most wellstudied model organisms, there is a relative paucity of information regarding D. melanogaster's natural history, ecology and behaviour, including habitat, food resources, and natural enemies (but see Reaume & Sokolowski, 2006; David & Capy, 1988; Turelli

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

& Hoffmann, 1991; Schmidt, et al. 2005; Fleury et al. 2004; Wilfert & Jiggins, 2014; Stephan & Li, 2006). While anti-predator behaviours are well studied as targets of selection in prey (Juliano & Gravel, 2002; Stoks, McPeek & Mitchell, 2003; Magurran et al. 1992), the genetic bases of such behaviours are seldom investigated. Given the range of genetic and genomic tools available for D. melanogaster, along with its complex behavioural repertoire and suitability for experimental evolution, understanding the antipredatory behaviours persisting in a natural population of the fruit fly brings us one step closer to deciphering the molecular mechanisms for anti-predator behaviours. Previous work has examined the effects of natural enemies on population and community structures of *Drosophila spp*. Worthen (1989) studied the effects of predation by rove beetles (staphylinids) on the coexistence of three mushroom-feeding *Drosophila* species, and Escalante & Benado (1990) showed that ant predators regulate population densities of wild D. starmeri (cactophillic fruit fly). In D. melanogaster per se, the role of parasites in influencing larval and adult behaviours has been extensively studied (Milan, Kacsoh & Schlenke, 2012; Kacsoh et al., 2013; Polak & Starmer, 1998). Despite this literature, we know little about the predators of D. melanogaster adults in the wild, nor the nature of anti-predatory behaviours segregating in natural populations. We documented the influence of two predators, the zebra jumping spider (Salticus scenicus) and juvenile Chinese praying mantids (Tenodera aridifolia sinensis) on the capture-deterrence behaviours of D. melanogaster individuals derived from a wild-caught population. The zebra spider is an active hunter, locating prey visually (with an extensive visual field attained by antero-medially positioned simple eyes) (Dill, 1975; Horner, Stangl & Fuller, 1988). Mantids are generally ambush predators, waiting for prey to enter

88

89

90

91

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

their attack range (Prete, Klimek & Grossman, 1990). Despite numerous differences, zebra spiders and juvenile Chinese mantids are similar in two relevant ways. First, both species primarily detect prey visually (Forster, 1979; Harland, Jackson, Macnab, 1999; Jackson, & Blest, 1982; Prete 1999) and are likely incapable of depth perception when their prey item is motionless (Prete, 1999; Freed, 1984). Second, small adult diptera account form a substantial proportion of the diet of both predators in the wild (Iwasaki, 1998; Okuyama, 2007). Based on the findings of Schmitz (2008), we predicted that fruit flies, in the presence of a familiar predator, would exhibit hunting-mode specific modifications in activity levels. To maximize distance from the actively hunting spider, our prediction was that flies would increase their overall activity levels, whereas, to reduce the probability of encountering a stationary threat (the mantid, an ambush predator), we expected flies to decrease overall activity. Under controlled laboratory conditions, we documented the behaviours of individual adult D. melanogaster with and without the two predator species. Our results suggest that in the presence of zebra spiders, D. melanogaster increases its overall locomotory activity, performs a distinct "stopping" behaviour and increases the performance of a newly described abdominal lifting behaviour (the function of which is as of yet unknown). Counter to our prediction though, D. melanogaster's locomotion, and most other behaviours are not substantially altered in the presence of mantids. However, upon direct encounter with a mantid, many individuals of D. melanogaster perform (a previously undescribed) retreat behaviour- a response not generally elicited by jumping spiders. Furthermore we observe considerable intra- and inter-individual variation in

response to predators. We discuss our results in terms of conditionally expressed behaviours as they relate to predator hunting mode, co-evolutionary history of predators and prey, and in terms of broadening our understanding of the behavioural ecology of D. melanogaster.

## **METHODS**

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

# Drosophila Population and Culture Conditions

The *Drosophila melanogaster* population used in this study originated from a natural population at Fenn Valley Vineyards in Fennville, Michigan (GPS coordinates: 42.57, -86.14) during the summer of 2010. A lab population (henceforth referred to as FVW) was initiated from this collection using the progeny of over 500 single-pair matings of field caught D. melanogaster as well as wild caught males. This design allowed us to screen out the sympatric congener, D. simulans, which was present in our collections at a frequency of about 5%. Screening involved setting up single pair mating in vials and discarding all lines with D. simulans-like genital morphology. After screening, ~1500 individuals were placed into cage (32.5cm<sup>3</sup>, BugDorm BD43030F) to establish the FVW population. The population is currently maintained in this cage at an adult density  $\sim 3000$  individuals in a room maintained at  $23^{\circ}$ C (+/-  $1^{\circ}$ C), and 40-70% RH. Adults were allowed to lay eggs in 10 bottles with 50-60 ml of a standard yeast-cornmeal food for 2-3 days. These bottles were then removed and kept in a Percival incubator (Model: I41VLC8) at 24<sup>o</sup>C and 65% RH throughout the larval stages. All flies and larvae were maintained in a 12 hr light/dark cycle with lights on at 08:00 hours.

133

134

135

136

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

For the experiments, pupae were collected 24 hours before they emerged as adults. Pupae were removed from bottles using forceps and individual pupae were placed into 1.5 ml microcentrifuge tubes. Each tube was pre-filled with  $\sim 0.5$  ml of yeastcornmeal food and its cap was punctured for gas exchange. Upon emergence, adult flies were sexed visually without anesthesia and housed in these tubes in the incubator until needed for behavioural assays. Age of flies used in behaviour analysis was 3-7 days. By using socially naïve flies in our assays, we were able to establish a consistent baseline of social experience among all individuals, allowing us to eliminate the potentially confounding influence of variation in social experience on behaviour that is welldocumented in *Drosophila* (Yurkovic et al., 2006; Levine, 2004; Krupp et al., 2008; Lefranc et al., 2001; Chabaud et al., 2009). **Spiders** S. scenicus individuals were collected throughout the spring/summer of 2012 on the campus of Michigan State University. Spiders were housed individually in vials in a room maintained at 23°C (+/- 1°C) and 30-50% RH and fed ~5 D. melanogaster a week. Prior to use in behavioural assays, spiders were starved for at least 48 hours. Each spider was used in only a single behavioural assay. Mantids Mantid egg cases were both collected near the campus of Michigan State University as well as ordered from Nature's Control (Medford, Oregon). Mantid egg cases were stored at 4°C and transferred to 25°C and 70% RH for hatching. Given the

substantial changes in mantid body size across moults (Iwasaki, 1990), only first instar nymphs were used for experiments. Prior to behavioural assay, mantids were starved for at least 24 hours and each mantid was used only once.

### **Behavioural** Assays

cues.

All assays were performed 1-4 hours after the incubator lights came on in the morning (08:00). Behavioural assays were recorded with an Aiptek AHD H23 digital camcorder attached to a tripod under a combination of natural and fluorescent light that is present in the room wherein the FVW population and spiders are maintained. For each predator (spiders and mantids), we recorded the behaviour for each of 15 male and 15 female socially naïve, virgin flies (collected as described above). We used a chamber constructed from the bottom of a 100 x15mm petri dish inverted on top of a glass plate with a sheet of white paper beneath to maximize the visibility of flies and predators.

For each assay, an individual fly was aspirated into the chamber and allowed to acclimate for 5 minutes. After this acclimation period, flies were recorded for 5 minutes. A single spider or mantid was then introduced to the chamber and behaviours were recorded for an additional 10 minutes or until capture. The chamber was washed with 10-30% ethanol and rinsed with reverse osmosis water after each assay to remove olfactory

### Behaviours Recorded

All *Drosophila* behaviours were categorized and analysed as either "states" or "events". Behavioural states have measurable duration and are mutually exclusive with

other states (e.g. individuals cannot simultaneously walk and run). Behavioural events are discrete behaviours that occur instantaneously and are also mutually exclusive with each other (e.g. turning versus jumping) but not always mutually exclusive with behavioural states. For example, an individual could perform a wing display (event) while simultaneously walking (state), but it could not jump (event) while simultaneously running (state). In this study we treated flying as an event because the structure of the experimental chamber constrained flight duration. Attempted flight by D. melanogaster could result in landing due to contact with a wall of the petri dish. We also recorded when a fly was not visible (occluded) to the observers analysing video. We recorded a total of 6 discrete events and 5 behavioural states in D. melanogaster in response to predation by spiders and mantids (Table 1). In order to interpret an individual fly's behaviour in the context of predatory encounters, we designated two keys to describe the location of the predator in regard to its interactions with the fly. As flies might alter their behaviour when a predator is within striking distance, we recorded predator location based on whether or not it was within striking distance of the fly (~5mm from the spider/mantid, also see *Spider location/ Mantid location* in Figure 1).

#### Video Processing

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

193

194

195

196

197

198

199

Recorded behaviours were viewed with VLC media player (version 2.0.3) and analysed by two observers using a manual event recorder, JWatcher V1.0 software (Blumstein, 2006). One observer (A.P.) viewed each video and verbally announced the occurrence of behaviours while the other observer (C.P./ M.C.) recorded the occurrence

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

221

222

of these behaviours with JWatcher. Because *Drosophila* anti-predatory behaviours are often complex and occur rapidly, we analysed all videos at 0.5X speed. Controlling for effects of season and disturbance We conducted experiments with spiders between October and December 2012 and those with mantids from March and May 2013. To confirm that predator species-specific behavioural differences were not confounded with seasonal differences in behaviour, we performed 6 additional assays (alternating between spider and mantid treatments) within the span of one week. Following a spider assay, the plates were wiped down with 30% ethanol followed by a rinse with RO water before a mantid assay was conducted. Additionally, the process of adding a predator to the arena invariably resulted in a disturbance that likely startled the fly (unrelated to the presence of a predator). To confirm that behaviours induced by this disturbance were not confounded with predator induced behavioural differences, we performed 3 control assays. Here, after 5 minutes of acclimatization without a predator (see above for more details), the arena containing the fruit fly was disturbed gently (~ magnitude of disturbance caused by the addition of a predator). For all controls, video processing and behaviours recorded were identical to mantid and spider treatments described above. See Supplement b, S1 for a detailed description of these control experiments and their results. Data processing and statistical analysis A custom Python script was used to parse Jwatcher formatted data files into a comma-separated-value (CSV) file for analysis in **R** (version 3.0.1).

To analyse the effects of predator state (i.e., presence or absence of predators) on the time dedicated to locomotory behavioural states, and number of occurrence for behavioural events, we fit generalized linear mixed effects models (using both <code>glmer</code> function in the lme4 package version 1.0-5, and the <code>MCMCglmm</code> function in the MCMCglmm package version 2.17) with predator state, total duration of assay with and without a predator (duration), sex, temperature and recording time as fixed effects, and individual by predator state and date as random effects . Formally, the model was:

$$y\sim\beta_0+\beta_1PS+\beta_2D+\beta_3Ag+\beta_4T+\beta_5ST+\beta_6Sx+\beta_7+\epsilon$$

Where  $\mathbf{y}$  is a vector of time spent in a behavioural state.  $\beta_1$  is the regression coefficient for predator state,  $\beta_2$  is for duration in each predator state,  $\beta_3$  is for age of the fly,  $\beta_4$  is for temperature,  $\beta_5$  is for time at which assay was started,  $\beta_6$  is for sex of the fly and  $\beta_7$  is for date on which the assay was performed. We estimated random effects for individuals including variation in response to predator state and duration of assay, and we fit an independent random effect for date. Thus we fit a repeated effects (longitudinal) mixed effects model allowing for variation among individuals for the influence of predator presence and duration of assay where for the  $i^{th}$  individual

$$\begin{pmatrix} \beta_{0i} \\ \beta_{1i} \\ \beta_{2i} \end{pmatrix} \sim MVN \begin{pmatrix} \begin{bmatrix} \mu_{\beta_0} \\ \mu_{\beta_1} \\ \mu_{\beta_2} \end{bmatrix}, \begin{bmatrix} \sigma_{\beta_0}^2 & \sigma_{\beta_0,\beta_1} & \sigma_{\beta_0,\beta_2} \\ \sigma_{\beta_0,\beta_1} & \sigma_{\beta_1}^2 & \sigma_{\beta_1,\beta_2} \\ \sigma_{\beta_0,\beta_2} & \sigma_{\beta_1,\beta_2} & \sigma_{\beta_2}^2 \end{bmatrix}$$

and (independent of the above)

244 
$$\beta_7 \sim N(0, \sigma_i^2)$$
 where  $i = 1 \dots$  date

246

247

248

249

250

251

252

253

254

255

256

257

258

259

260

261

262

263

264

265

266

267

Preliminary analyses were inconsistent with the need to fit higher order interactions among fixed effects, so interaction terms were not considered further. The one exception was for "stopping" behaviour where individuals almost exclusively performed this in the presence of the predators. For the behavioural states (locomotion, grooming and stopping) we assumed normally distributed variation. For the counts of events (abdominal lift, jumping, etc) we used a log-link function and assumed the variation was poisson distributed. Estimation using both maximum likelihood (lmer) and simulating the posterior distribution (MCMCqlmm) provided similar results for fixed effects, and generally for random effect components as well. Among individual coefficients of variation were calculated by dividing the square root of the among individual variance component from the model by its respective fixed effect estimate (i.e. its "mean"). While confidence intervals were consistent for fixed effects, the intervals were more difficult to estimate given the complexities of the random effect structure of the model, and some caution is warranted for their interpretation. To test for non-random associations in the temporal structure of behavioural patterns we constructed transition frequencies using the "msm" library (version 1.2) (Jackson, 2011) in **R**. To test for both for first order Markov processes between behaviours (transition probabilities), as well as the influence of predator presence on these transition probabilities, we fit log-linear models (assuming poisson distributed data) with the transition frequency matrices (Crawley, 2012) using alm in R. As advocated by (Crawley, 2012; Bakeman & Gottman, 1997) we fit a saturated log-linear model (with lag0, lag1 and predator state as the effects in the model) and tested the influence of

# Table 1 Names and descriptions of all observed behaviours. Videos

are provided at the end of Supplement b.

Behaviour	Description
Abdominal lift (ab)	Momentary rearing up on abdomen (see video 1)
Fly	Moving through space by wing use
Jump	Instantaneous movement between points without wing use
Pause	Noticeable period of inactivity; transitional
Turn	180 degree change in orientation without change in position
Wing display (wd)	Momentary lifting up of wings without singing or vibration
Groom	Running legs over any body part-often while otherwise stationary
Walk	Movement through space by ambulation
Run	Rapid movement through space by ambulation
Stop	Immobile (see video 2)
Retreat	Walking in reverse upon encounter with an object (like a predator) (see video 3)

269

270

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

286

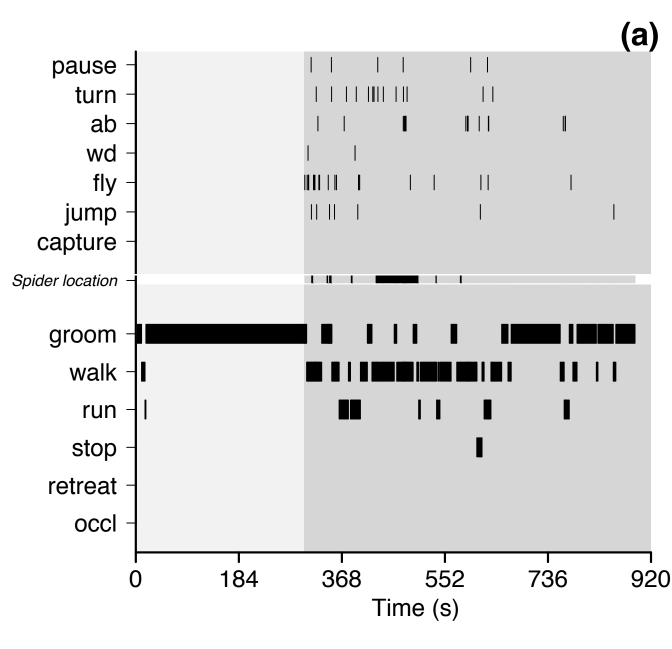
287

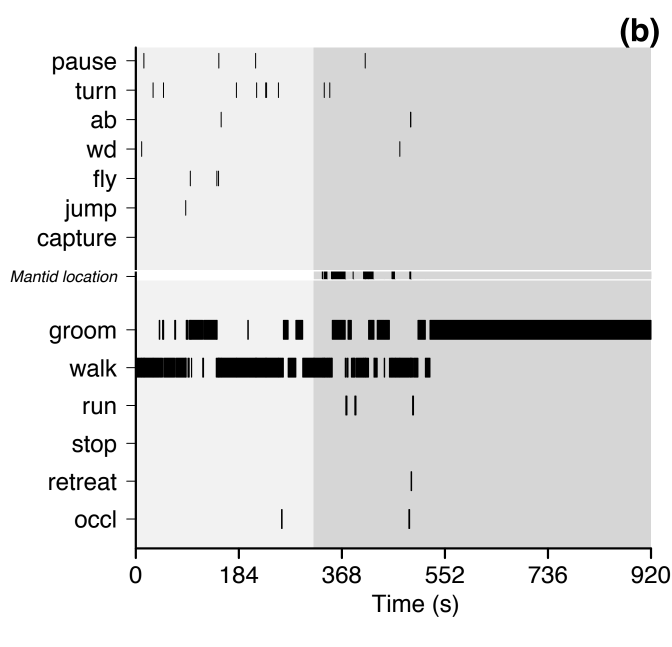
288

289

290

deleting the terms (i.e. third order interaction) on change in deviance. We used modified "Z-scores", adjusted using sequential Bonferroni to assess the deviation of particular cells in the transition frequency matrix from expected values (assuming independence). For the visual transition probability matrices, we combined the behavioural event "pause" with the behavioural state "stop" because 1) we wanted to reduce the complexity of the matrix and 2) the main difference between the two behaviours is that pause is instantaneous and stop has duration. All transition diagrams were constructed in Inkscape (version 0.48.2, Harrington, 2004-2005). **RESULTS** From pilot observations (not included in analysis), we (I.D., A.P. and C.P.) catalogued and described Drosophila melanogaster behaviours observed in the presence of a predator (Table 1). Among the behaviours listed in Table 1, abdominal lifting (ab, supplement b, video 1) and retreat (supplement b, video 3), to our knowledge, have not been previously described in *D. melanogaster* literature. Flies perform a range of anti-predatory behaviours in response to a zebra spider To visualize each individual fruit fly's response to the presence of a zebra jumping spider, we generated ethograms (see Figure 1a and Supplement a). For the two predator states (spider present and spider absent) we measured the mean proportion of time dedicated to each behavioural state, as well as the number of occurrences per minute for each behavioural event. When a spider was present, D. melanogaster increased the proportion of time it spent walking and running by 50% (95% CI: 21-79% increase)





# Figure 1

D. melanogaster used a greater proportion of its behavioural repertoire and performed each behaviour at a higher frequency in the presence of a jumping spider than in the presence of a juvenile mantid. a) Representative ethogram of a male, 4 day old *D. melanogaster* in response to a zebra jumping spider. b) Ethogram of a male, 5 day old D. melanogaster in response to a juvenile Chinese paying mantid. Light grey background represents time in the arena before the addition of a predator and dark grey background is when the predator was present in the chamber. Each black bar represents the occurrence of a behaviour during the observation period. Top half of the figure (separated by *Predator location*) consists of events and the bottom half consists of states. Because states have duration, the width of each black bar corresponds to the duration of a state. Predator location (i.e., Spider location in a and Mantid location in b) indicates whether or not the predator was within striking distance of the fruit fly at that time point. This information is relevant only after the predator was added to the chamber (~ 300 s into the assay). Dark grey bars in *Predator location* indicate that the spider was within striking distance and light grey regions indicate that the spider was out of striking distance. *Predator* location is white when the predator is absent from the arena or after successful capture. If capture did not occur, predator location remains light grey in colour.

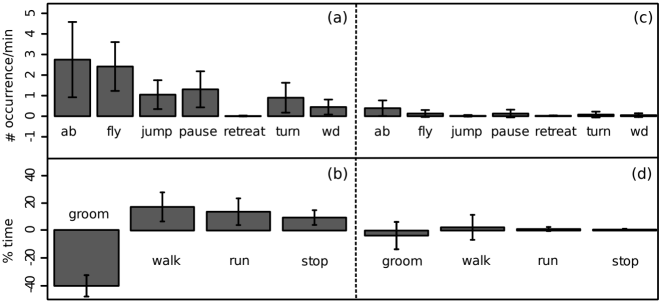


Figure 2

Fruit flies increase overall activity levels in the presence of jumping spiders (a) and (b) but not in the presence of mantids (c) and (d). Plots (a) and (b) show change in mean number of occurrences per minute of each behavioural state as a result of the addition of a predator. Plots (c) and (d) show mean change in percentage of total time spent in a given behavioural state caused by the addition of a predator. On the left of the dotted line, behavioural changes correspond to the presence of a spider whereas on the right of the dotted line, behavioural differences are due to the presence of a juvenile praying mantid. Error bars are  $\pm$  95% CI

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

312

313

while grooming 60% less (95% CI: 43-77% decrease). This is shown in Figure 2 and Supplement b Figure S1 (treatment contrasts with 95% CI in figures are provided to enable assessment of significance). While they were observed at low frequencies prior to the addition of spiders, D. melanogaster substantially increased the frequency of pauses, jumps and flights (per minute) in the presence of spiders (Figure 2, Supplement b Figure S3). For instance the frequency of abdominal lifts increased from 1.51/minute to 4.0/minute (95% CI: 1.55-9.93), while jumping showed a 6.6X increase from 0.73/minute to 4.82/minute (95% CI: 1.77-12.17). "Stopping"; a motionless state that likely aids in capture deterrence (see Videos 2, Supplement b) was not performed by D. melanogaster in the absence of spiders (Supplement b Figure S2). However in the presence of spiders, the average total time spent "stopping" increased to ~25.8 seconds (95% CI: 10.1 - 41.7 seconds). When interacting with spiders, flies were only observed to perform the "retreat" behaviour once (of 30 individuals). Interestingly, we did not see significant sex specific differences in either frequencies of occurrence (Supplement b Figure S3) or proportion of time allocate (Supplement b Figure S1) to the majority of measured behaviours (But see S3 panels "pause" and "turn"). Given the design of our experiment, we were able to model the degree to which individuals varied in their responses to the jumping spiders. Individuals varied greatly both in their baseline activity levels as well as in their propensities to respond to jumping spiders. The among-individual coefficient of variation for time spent grooming in the absence of predators was 57.7% (40.1-74.2%). While most individuals reduced their grooming activity in the presence of predators, the degree to which they did so varied substantially, with the among individual coefficient of variation for the decrease being

315

316

317

318

319

320

321

322

323

324

325

326

327

328

329

330

331

332

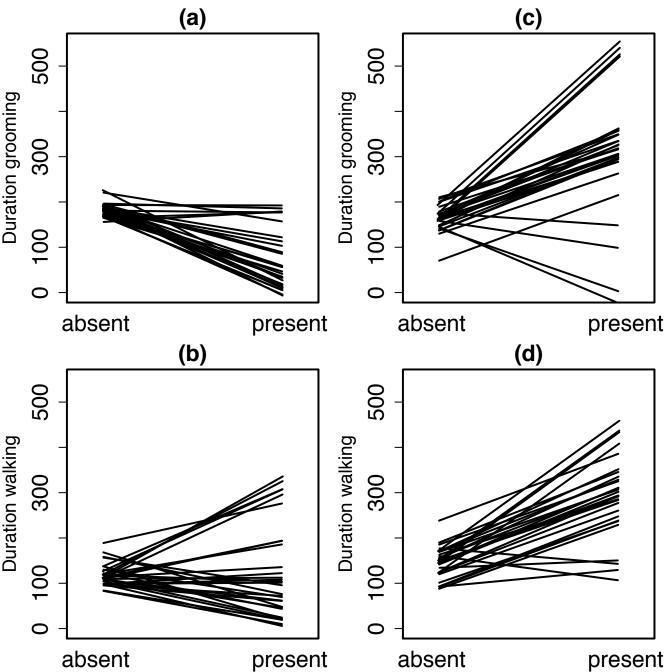
333

334

335

336

67.8% (26.5-94.6%), as shown in Figure 3a. For walking, the among-individual coefficient of variation was 80.3% (50-105%) in the absence of the spider, and 135% (1-181%) for the magnitude of increase in the presence of the spider (Figure 3b). Performance of the stopping behaviour by D. melanogaster in the presence of spiders varied substantially among individuals, with the among-individual coefficient of variation being 168% (95% CI: 123-214%). This is driven in part by the fact that 40% of individual flies never performed stopping, even in the presence of the spider. There was a negative correlation (-0.84), between the amount of time individuals spent grooming before and after the addition of the spiders (Table 2). That is, on average, individuals who were more active prior to the addition of the spider reduced their activity to a greater extent in the presence of the spider. A similar negative correlation (-0.66) for among individual activity for locomotion, was observed (Table 2). To visualise the temporal associations among behavioural sequences, we constructed transition matrices (Supplement b Tables S1, S2, S5 and S6) and transition probability diagrams for all pairs of behaviours in the presence (Figure 4a) and absence (Supplement b Figure S7) of spiders. In response to jumping spiders, transitions among behaviours are somewhat more dispersed (with many connections between behaviours), suggesting that there is weak temporal association between fruit fly behaviours. Indeed these qualitative conclusions are supported based on the Z-scores. In the absence of spiders 8 possible transitions were significant (after controlling for multiple comparison, Supplement b Table S2), while 13 transitions were significant in the presence of the spider (Supplement b Table S1). Most of these differences were due to the increase in behaviours potentially involved with anti-predation activity (i.e. flight, abdominal



# Figure 3

models.

Inter individual behavioural variation in response to predators is present in natural populations. Reaction norms visualize how each individual fruit fly responded to the introduction of a spider (panels a and b) or a mantid (panels c and d) into the assay chamber. Measures are in seconds. Each line corresponds to response of one individual. Estimates are derived from the predicted values for each individual from the mixed

Table 2 Individual flies show negative correlations between behavioural states before and after the introduction of a predator. There is considerable variation among individuals in time spent performing specific behaviours (i.e. walking and grooming), with and without predators. However, there is a strong negative correlation within individuals for time spent before and after introduction of the predator. That is, individuals who spend more time performing a specific behaviour prior to the addition of a predator, reduce that behaviour to an even greater amount (than the average for the sample) once the predator is introduced. The one exception is for grooming for the mantid trials. Diagonals of the table contain the standard devation (mean of the posterior distribution) for individual behavioural responses (95% CIs in paratheses) from the random effects of the models. Above the diagonal are covariances between predictors (and CIs in parantheses). Below the diagonal are correlation coefficients for the covariances between the predictors.

**Grooming, Spider Pred.State** Intercept Time 89.2 (62.0, 115.3) 26.4 (14.0, 36.0) Intercept -68.2 (-88.7, -34.1) **Pred.State** -0.8461.7 (18.8, 84.5) -14 (-25.5, 17.1) 0.75 -0.310.4 (4.4, 14.9) Time Walking, Spider **Pred.State** Time Intercept 81.9 (43.3, 109.8) -60.2 (-89.5, 27.5) 25.6 (-8.8, 36.7) Intercept Pred.State -0.6667.3 (0.36, 98.5) 11.5 (-22.2, 29.2) **Time** 0.62 0.15 13 (5.6, 18.2) **Grooming, Mantid** Intercept Pred.State Time Intercept 122.8 (59.6, 175) -20.2 (-117, 106) 46.0 (-16.9, 74.5) **Pred.State** -0.05 60.5 (0.13, 100.2) -18.3 (-47.9, 34.0) 8.0 -0.2621.5 (2.6, 33.8) Time Walking, Mantid Intercept Pred.State Time Intercept 144.8 (86.2, 198) -100.3 (-162.6, 38.8) 63.2 (31.6, 90.3) Pred.State -0.8680.5 (0.21, 139.3) -45.2 (-76.4, 19.6) 0.94 -0.8629.4 (11.3, 43.5) Time

338

339

340

341

342

343

344

345

346

347

348

349

350

351

352

353

354

355

356

357

358

lifting). However, while the results of the log-linear analysis (across the whole transition frequency matrix) supported the dependence of current behavioural states on the previous state (resid df=71, deviance=632, p < 0.001), the inclusion of predator status did not influence this dependence (resid df = 71, deviance = 59, p = 0.8). Flies perform a previously undescribed retreat behaviour in response to mantids In contrast to their behaviour in the presence of jumping spiders, the presence of a juvenile mantid had a minimal influence on D. melanogaster's locomotory activity (Figure 2c and 2d, also see ethograms in Figure 1b and Supplement a). Time spent grooming, walking, running and stopping was highly variable (with estimates including zero) in the presence of a juvenile praying mantid (Figure 2d, Supplement b Figures S1 and S2). Grooming decreased by 20% (95% CI: 83% decrease to 36% increase), locomotion increased by 9% (13% decrease to 32% increase) and stopping increased by 40% (8% decrease to 85% increase). Similarly, the presence of a mantid did not influence the frequency at which D. melanogaster tended to perform most instantaneous behaviours (Figure 2c, Supplement b Figure S4). However, as was observed in the presence of spiders, flies substantially increased the frequency of abdominal lifting, from 0.45/minute to 5.6/min (95% CI: 2.56 - 12.17) in the presence of a juvenile mantid (Supplement b Figure S4). Upon encounter with a mantid, half of the individuals (15/30) performed a previously undescribed reversal behaviour (Supplement b video 3), which we term "retreat". As with the zebra spiders, we saw no significant sex specific differences in response to mantids.

360

361

362

363

364

365

366

367

368

369

370

371

372

373

374

375

376

377

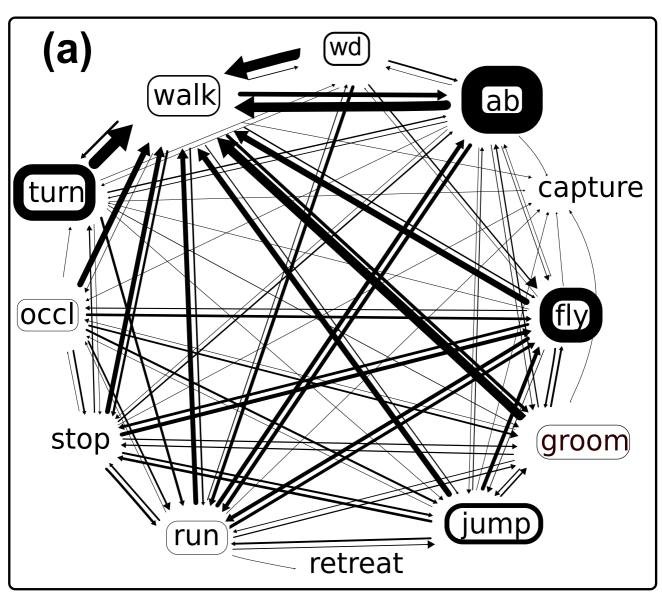
378

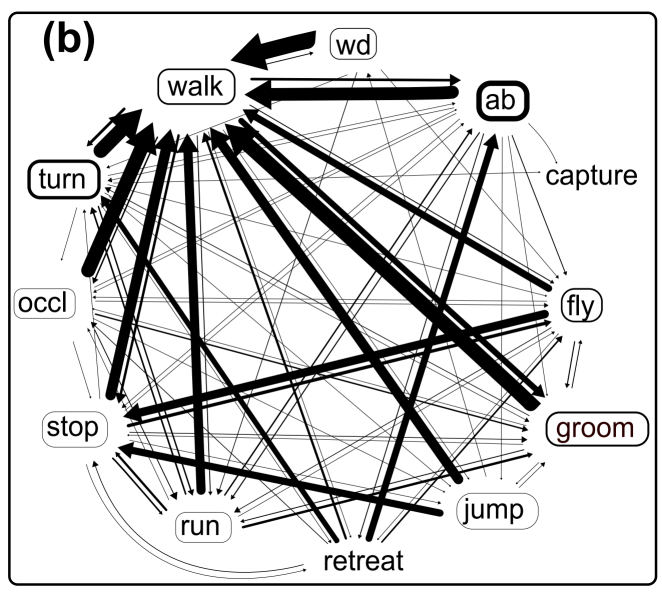
379

380

381

Although the presence of a mantid had a small effect on fly behaviour, flies did vary considerably in their grooming and walking activities. Indeed, the among-individual variability in proportion of time spent grooming and walking is greater in magnitude in the presence of the mantids than spiders (Figure 3). Evidence for negative co-variation for intra-individual behaviour before and after the addition of the predator was not strongly supported (i.e. 95% CIs for covariances included zero) (Table 2). Transition matrices and transition probability diagrams (Supplement b Figure S7b, Figure 4b and Tables S3, S4, S7 and S8) show patterns of temporal association among behaviours. In response to juvenile mantids, the transitions diagram is less dispersed than that in the presence of jumping spiders (Figure 4), suggesting that the degree of association between behaviours in the presence of mantids is more extreme. While most behaviours (abdominal lift, fly, groom, jump, run, stop, and turn) tend to transition to walking, we also see stronger associations between other pairs of behaviours. For example, after performing the retreat behaviour, flies often either performed the abdominal lift or turn, while flight is often followed by stopping. These observations are supported by the findings that in the absence of mantids, 12 transitions showed significant deviations from expectations (Supplement b, Table S4). In comparison, in the presence of mantids 23 transitions showed a significant deviation from expected values (Supplement b, Table S3). Interestingly, as with the spiders the log-linear model supports the non-independence of behavioural states (resid df=71, deviance=1054, p <0.001), but not for the additional influence of predator state on this non-independence (resid df = 71, deviance = 72, p=0.4).





# Figure 4

Spiders and mantids had different effects on the temporal associations between pairs of D. melanogaster behaviours. a) A diagram representing probability of transitioning from one fly behavioural state to the other in the presence of a zebra jumping spider. b) A diagram representing probability of transitioning from one fly behavioural state to the other in the presence of a juvenile praying mantid. Thickness of arrows indicates transition probability between the two behaviours. The arrowhead points to the behaviour being transitioned to. Thickness of the box around behavioural state (groom, run, occl, retreat, stop and walk) indicate the mean proportion of total time spent in that behaviour, whereas thickness of the box around behavioural events (fly, jump, turn, wd, ab) indicates mean number of occurrences per minute of that behaviour. To reduce the complexity of the web we combined the behaviours "pause" with the behaviour "stop". Behavioural transitions that occurred less than 10 times have not been shown in the figure.

### **DISCUSSION**

382

383

384

385

386

387

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

Prey organisms can alter their behaviour to reduce the likelihood of detection, capture or encounter with a predator (Lima, 1998). For example, when predators are present, ground squirrels dedicate more time to vigilance behaviours (like scanning for a predator, see (Bachman, 1993) and some aquatic insects spend more time in refuges (Kohler & McPeek, 1989). These changes in behaviour may alter the use of resources, and potentially the fitness of an organism. However, the nature and intensity of nonconsumptive effects of a predator on its prey are a function of several predator specific factors, one of which is the predator's hunting mode (Preisser, Orrock & Schmitz, 2007). Predator hunting mode, i.e., the set of behavioural strategies that a predator employs to pursue and capture its prey (Schoener, 1971; Huey & Pianka, 1981; Schmitz, 2008) can be an important determinant of a prey organism's anti-predatory behavioural response (Schmitz, 2008). In this study, we describe the anti-predatory behavioural repertoire of a natural population of *Drosophila melanogaster* in response to predation by the zebra jumping spider (Salticus scenicus) and juvenile Chinese praying mantids (Tenodera aridifolia sinensis). Among other characteristics, zebra spiders and praying mantids differ in their hunting mode. While we discuss our findings with respect to hunting mode differences, we recognize that other attributes differing among the predators may contribute to the observed differences in prey behavioural repertoires. However, as our experimental design was meant to minimize the effects of many possible confounding factors (e.g. time of day, temperature, humidity) it seems likely that, in part, our results reflect hunting mode differences.

405

406

407

408

409

410

411

412

413

414

415

416

417

418

419

420

421

422

423

424

425

426

In response to active hunters (those that constantly patrol for prey), we predicted that fruit flies would increase their overall activity levels (including flight) in order to maintain maximum distance from the predator at all times; To reduce the likelihood of an encounter with an ambush predator however (i.e., a predator that only attacks when a prey organism wanders in to its strike zone), we predicted that D. melanogaster would respond by decreasing locomotory activities (Schmitz, 2008). Our results, however, were only partially in line with these predictions. While the actively hunting jumping spiders induce a clear increase in overall activity, we found the presence of juvenile mantids- our ambush predators- to have minimal influence on fruit fly activity levels (Figure 2, Supplement b Figure S2). It has been previously argued that ambush predators might be a predictable source of threat to prey organisms (Preisser, Orrock & Schmitz, 2007; Schmitz, 2008) as opposed to the diffuse and variable threat imposed by active hunters (Schmitz, 2008). Therefore, it is perhaps surprising that fruit flies show a stronger behavioural response to the threat of active hunters (zebra jumping spiders). However, our predictions are based on studies on a grasshopper and its two predatory spider species that differ in hunting mode. Given that selection pressures faced by adult diptera are different from those experienced by grasshoppers (orthoptera), such predictions may not be generalizable. Several factors including body size and dispersal patterns may contribute to this difference. Many species of jumping spiders, including S. scenicus, are often seen in the natural habitat of D. melanogaster (personal observations of A.P., C.P. and I.D.), and are likely to be ecologically relevant predators of *Drosophila*. Mantids however, are rarely found in areas where fruit flies are abundant (personal observations of A.P. and I.D.), at least in Eastern North America. Therefore, it is likely that fruit flies,

428

429

430

431

432

433

434

435

436

437

438

439

440

441

442

443

444

445

446

447

448

449

having experienced a longer evolutionary history with small jumping spiders, are better able to recognize these spiders as a threat. In addition, the disturbance created by a constantly patrolling zebra spider may be partly responsible for the increased activity levels seen in D. melanogaster (either due to actual mechanical disturbance or because flies are able to detect moving objects quicker than stationary ones). In this study, we are unable to tease apart the effects of evolutionary recognition versus constant mechanical disturbance on the differences in flies' activity levels. Further experimentation with harmless but constantly moving heterospecifics (such as field crickets) or immobilized active hunters might be useful in addressing these issues. We also identified a number of (to our knowledge) undescribed behaviours of D. melanogaster, potentially relating to its interactions with predators. The behaviour we called "stopping" (Table 1) was observed numerous times after a direct (but failed) attack by a spider (Supplement 3 video 1). While D. melanogaster will spend time without any ambulatory activity (walking, running), they are almost always observed to be active (generally grooming) during these periods. However, when fruit flies performed the stopping behaviour, there was a complete lack of movement on the part of the fly, even when video was viewed at a few frames/second. When a fruit fly was "stopped", the spider had to search for the fly, irrespective of the physical proximity between the spider and the fly. In salticids, while the principal eyes have high spatial acuity, secondary eyes are primarily used to detect moving objects (Harland, Jackson & Macnab, 1999; Land, 1971). Because salticids are unable to accommodate by changing the shape of their lens, they need to extensively sample their visual field to see details in object shape and form (Harland, Jackson & Macnab, 1999; Land, 1971; Blest, Hardie & McIntyre, 1981).

451

452

453

454

455

456

457

458

459

460

461

462

463

464

465

466

467

468

469

470

471

472

Scanning for prey by such sampling is likely a slow process unless guided by the motion sensing peripheral eyes, giving motionless prey the advantage of staying hidden (at least for a few seconds) while in plain sight of their salticid predator. Thus, D. melanogaster may be using the "stopping" behaviour as a potential mechanism to reduce the likelihood of detection by the spider. Additionally, in the presence of both predators, D. melanogaster substantially increase the frequency at which it performed abdominal lifts. To our knowledge, abdominal lifting has not been described in D. melanogaster literature before and may be relevant in an anti-predatory context. While studying courtship behaviours in female D. melanogaster, Lasbleiz, Ferveur & Everaerts (2006) described two behaviours perhaps similar to the abdominal lifting described here: abdominal drumming and abdominal extension. Abdominal drumming (described as "quickly repeated vertical movements of the abdomen which is tapped on the substrate") was only seen in males during courtship display, and abdominal extensions (described as "abdomen raised by 15-30 degrees") were also closely associated with courtship. Because abdominal lifting was often directed at a predator or followed a failed predatory encounter, we suspect abdominal lifting to be different from abdominal extensions and abdominal drumming, and with a possibly antipredatory function. We speculate that if abdominal lifting is indeed anti-predatory, it could function in one of several possible ways. First, abdominal lifting may be a signal of prey condition directed at the predator as a form of pursuit deterrence, comparable to stotting in the Thomson's gazelle (FitzGibbon & Fanshawe, 1988). Second, because D. *melanogaster* are often surrounded by conspecifics, abdominal lifting may be a means though which one fly warns its conspecifics of the presence of a potential threat (similar

474

475

476

477

478

479

480

481

482

483

484

485

486

487

488

489

490

491

492

493

494

in function to fin flicking in tetras, (Brown, Godin & Pedersen, 1999). Finally it may be an indication of some sort of physiological priming of the fly in preparation for a fightor-flight response. Determining whether it is a specific anti-predator behaviour, as well as the details of its function need to be a focus of future work. In response to the juvenile praying mantids, half of the fruit flies we observed (15/30) performed a reverse walking behaviour which we have called "retreat", where the flies walked in reverse, away from the predator (supplement b, video 3). This was often (but not always) interspersed with the abdominal lifting behaviour. Phenomenologically, this behaviour may be similar to that described in Bidaye et al. (2014). Bidaye et al. (2014) identified neurons that upon activation changed walking direction in D. melanogaster. Bidaye et al's reverse walking behaviour appears to be a smooth and continuous behaviour, whereas "retreat" was often discontinuous and accompanied by abdominal lifting. If the two "retreat" behaviours are related, the observed disassociation between retreat and abdominal lifting as well as its continuous nature (in Bidaye el al., 2014) may be a function of how the neurons were perturbed. We also investigated how the presence of the different predators may influence non-random associations among behaviours. We observed that in the presence of both predators there was an increase in the number of behavioural transitions that deviated from expectations under independence (from 12 to 23 with the mantid, and 8 to 13 with the spider). Despite this, the log-linear model (analysing the whole transition frequency matrix) did not support the influence of predator state on the frequencies of transitions. This may be partly due to the relatively modest sample sizes (in terms of both individuals

and transitions among behaviours). Further work is necessary to validate and extend this sequential analysis.

While we show that there are some predator hunting-mode specific behavioural differences in D. melanogaster's anti-predator response, we reiterate two important caveats. First, although the primary distinction between the zebra jumping spider and juvenile Chinese praying mantids as predators is their hunting-mode, other factors between these species (for example, size, colour, odour) may also influence differences in fruit fly behaviours. Replicating the observations with other predator pairs that differ in hunting-mode is necessary to confirm hunting-mode's influence on anti-predatory repertoires. Secondly, our assay chambers are an artificial environment and do not resemble the conditions under which D. melanogaster face predators in the wild. Due to the nature of our assay chamber, D. melanogaster were unable to employ behavioural strategies that may reduce encounters with predators (e.g., utilizing a refuge). Therefore we were only able to describe the capture-deterrence repertoire of D. melanogaster behaviour. We believe that our study is a necessary first step to describing and documenting the complete anti-predatory behavioural repertoire of D. melanogaster and we foresee future work to be conducted in a modified chamber, under more "natural" conditions. Doing so will allow us to take this premier model genetic system and make it into an ecological model as well.

## **ACKNOWLEDGEMENTS**

495

496

497

498

499

500

501

502

503

504

505

506

507

508

509

510

511

512

513

514

515

516

517

We thank Dr. Fred Dyer for loaning us video cameras. Thanks to Dr. Ben DeBivort for discussions and suggestions and to Dr. Reuven Dukas and Dr. Marla Sokolowski and

members of the lab for comments on the manuscript. This material is based in part upon work supported by the National Science Foundation under Cooperative Agreement No. DBI-0939454. Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the National Science Foundation.

524

525

526

527

528

529

530

531

532

533

534

535

536

537

538

539

540

541

542

REFERENCES Bachman, G. C. (1993) The effect of body condition on the trade-off between vigilance and foraging in Belding's ground squirrels. Anim Behav 46: 233-244. Bakeman, R. & Gottman, J. M. (1997) Observing interactions. New York: Cambridge University Press. Bidaye, S. S., Machacek, C., Wu, Y. & Dickson, B. J. (2014) Neuronal Control of Drosophila Walking Direction. Science 344, 97–101. (doi:10.1126/science.1249964) Blest, A. D., Hardie, R. C. & McIntyre, P. (1981) The spectral sensitivities of identified receptors and the function of retinal tiering in the principal eyes of a jumping spider. J. Comp. Physiol., A 145: 227-239. Blumstein, D. T. (2006) The multi-predator hypothesis and the evolutionary persistence of antipredator behavior. Ethology 112: 209–217. (doi:10.1111/j.1439-0310.2006.01209.x) Brown, G., Godin, J. & Pedersen, J. (1999) Fin-flicking behaviour: a visual antipredator alarm signal in a characin fish, *Hemigrammus erythrozonus*. Anim Behav 58: 469–475. (doi:10.1006/anbe.1999.1173) Chabaud, M.A., Isabel, G., Kaiser, L. & Preat, T. (2009) Social facilitation of longlasting memory retrieval in *Drosophila*. Curr Biol 19: 1654–1659. (doi:10.1016/j.cub.2009.08.017) Crawley, M. J. (2012) The R book. John Wiley & Sons.

543 David, J. R. & Capy, P. (1988) Genetic variation of *Drosophila melanogaster* natural 544 populations. Trends Genet 4: 106–111. 545 DeWitt, T. J., Robinson, B. W. & Wilson, D. S. (2000) Functional diversity among 546 predators of a freshwater snail imposes an adaptive trade-off for shell morphology. Evol 547 Ecol Res 2: 129–148. 548 Dill, L. M. (1975) Predatory behavior of the zebra spider, Salticus scenicus (Araneae: 549 Salticidae). Can J Zool 53: 1284–1289. 550 Edmunds, M. (1974) Defence in animals: a survey of anti-predator defences. Longman Harlow, New York. 552 Escalante, A. & Benado, M. (1990) Predation on the cactophilic fly, *Drosophila starmeri*, 553 in the columnar cactus, *Pilosocereus lanuginosus*. Biotropica, 48–50. 554 FitzGibbon, C. D. & Fanshawe, J. H. (1988) Stotting in Thomson's gazelles: an honest 555 signal of condition. Behav Ecol Sociobiol 23: 69–74. 556 Fleury, F., Ris, N., Allemand, R., Fouillet, P., Carton, Y. & Boulétreau, M. (2004) 557 Ecological and genetic interactions in Drosophila-parasitoids communities: a case study 558 with D. melanogaster, D. simulans and their common Leptopilina parasitoids in south-559 eastern France. Genetica 120: 181–194. 560 Forster, L. M. (1979) Visual mechanisms of hunting behaviour in *Trite planiceps*, a 561 jumping spider (Araneae: Salticidae). NZ J Zool 6: 79–93. 562 (doi:10.1080/03014223.1979.10428351)

564

565

566

567

568

569

570

571

572

573

574

575

576

577

578

579

580

581

582

Freed, A. N. (1984) Foraging behaviour in the jumping spider *Phidippus audax*: bases for selectivity. J Zool 203: 49-61. Gosline, A. K. & Rodd, F. H. (2007) Predator-induced plasticity in guppy (*Poecilia* reticulata) life history traits. Aquat Ecol 42, 693–699. (doi:10.1007/s10452-007-9138-7) Harland, D. P., Jackson, R. R. & Macnab, A. M. (1999) Distances at which jumping spiders (Araneae: Salticidae) distinguish between prey and conspecific rivals. J Zool 24: 357-364. Harrington, B. et al (2004-2005) Inkscape. http://www.inkscape.org/ Horner, N. V., Stangl, F. B. & Fuller, G. K. (1988) Natural history observations of Salticus austinensis in north-central Texas. J Arachnol 16: 260-262. Huey, R. B. & Pianka, E. R. (1981) Ecological consequences of foraging mode. Ecology 4: 991-999. Iwasaki, T. (1990) Predatory behavior of the praying mantis, *Tenodera aridifolia* I. Effect of prey size on prey recognition. J Ethol 8: 75–79. Iwasaki, T. 1998 Prey menus of two praying mantises, Tenodera aridifolia (Stoll) and Tenodera angustipennis Saussure (Mantodea: Mantidae). Entomol Sci 4: 529-532. Jackson, R. R. & Blest, A. D. (1982) Short Communication: The distances at which a primitive jumping spider, *Portia fimbriata*, makes visual discriminations. J Exp Biol 97: 441–445. Jackson, C. H. (2011) Multi-State Models for Panel Data: The msm Package for R. J Stat

583 Softw 38(8): 1-29. URL http://www.jstatsoft.org/v38/i08/ 584 Juliano, S. A. & Gravel, M. E. (2002) Predation and the evolution of prey behavior: an 585 experiment with tree hole mosquitoes. Behav Ecol 13, 301–311. 586 Kacsoh, B. Z., Lynch, Z. R., Mortimer, N. T. & Schlenke, T. A. (2013) Fruit Flies 587 Medicate Offspring After Seeing Parasites. Science 339: 947–950. 588 (doi:10.1126/science.1229625) 589 Kohler, S. L. & McPeek, M. A. (1989) Predation risk and the foraging behavior of 590 competing stream insects. Ecology 70: 1811–1825. 591 Krupp, J. J., Kent, C., Billeter, J.-C., Azanchi, R., So, A. K. C., Schonfeld, J. A., Smith, 592 B. P., Lucas, C. & Levine, J. D. (2008) Social experience modifies pheromone expression 593 and mating behavior in male *Drosophila melanogaster*. Curr Biol 18: 1373–1383. 594 (doi:10.1016/j.cub.2008.07.089) 595 Langerhans, R. B. (2007) Evolutionary consequences of predation: avoidance, escape, 596 reproduction, and diversification. Predation in organisms 177-220. Springer Berlin 597 Heidelberg. (doi: 10.1007/978-3-540-46046-6 10) 598 Land, M. F. (1971) Orientation by jumping spiders in the absence of visual feedback. J 599 Exp Biol 54: 119–139. 600 Lasbleiz, C., Ferveur, J.-F. & Everaerts, C. (2006) Courtship behaviour of *Drosophila* 601 melanogaster revisited. Anim Behav 72: 1001–1012.

602

(doi:10.1016/j.anbehav.2006.01.027)

603 Lefranc, A., Jeune, B., Thomas-Orillard, M. & Danchin, E. (2001) Non-independence of 604 individuals in a population of *Drosophila melanogaster*: effects on spatial distribution 605 and dispersal. C R Acad Sci III, Sci Vie 324: 219–227. 606 Levine, J. D. (2004) Sharing time on the fly. Curr Opin Cell Biol 16: 210–216. 607 (doi:10.1016/j.ceb.2004.02.009) 608 Lima, S. L. & Dill, L. M. (1990) Behavioral decisions made under the risk of predation: a 609 review and prospectus. Can J Zool 68, 619–640. 610 Lima, S. L. (1998) Nonlethal effects in the ecology of predator-prey interactions. 611 BioScience 48, 25–34. 612 Lima, S. L. & Bednekoff, P. A. (1999) Temporal variation in danger drives antipredator 613 behavior: the predation risk allocation hypothesis. Am Nat 153: 649-659 614 Magurran, A. E., Seghers, B. H., Carvalho, G. R. & Shaw, P. W. (1992) Behavioural 615 consequences of an artificial introduction of guppies (*Poecilia reticulata*) in N. Trinidad: 616 Evidence for the evolution of anti-predator behaviour in the wild. Proc R Soc B 248: 617 117–122. (doi:10.1098/rspb.1992.0050) 618 Milan, N. F., Kacsoh, B. Z. & Schlenke, T. A. (2012) Alcohol consumption as self-619 medication against blood-borne parasites in the fruit fly. Curr Biol 22: 488–493. 620 (doi:10.1016/j.cub.2012.01.045) 621 Okuyama, T. (2007) Prey of two species of jumping spiders in the field. Appl Entomol 622 Zool 42: 663–668. (doi:10.1303/aez.2007.663)

623 Polak, M. & Starmer, W. T. (1998) Parasite-induced risk of mortality elevates 624 reproductive effort in male *Drosophila*. Proc Royal Soc B 265: 2197–2201. 625 Preisser, E. L., Orrock, J. L. & Schmitz, O. J. (2007) Predator hunting mode and habitat 626 domain alter nonconsumptive effects in predator-prey interactions. Ecology 88: 2744– 627 2751. 628 Prete, F. R., Klimek, C. A. & Grossman, S. P. (1990) The predatory strike of the praying 629 mantis, Tenodera aridifolia sinensis. J Insect Physiol 36: 561-565. 630 Prete, F. R. (1999) The Praying Mantids 93-141. Baltimore: The Johns Hopkins 631 University Press. 632 Reaume, C. J. & Sokolowski, M. B. (2006) The nature of *Drosophila melanogaster*. Curr 633 Biol 16: R623-R628. 634 Relyea, R. A. (2001) Morphological and behavioral plasticity of larval anurans in 635 response to different predators. Ecology 82: 523–540. 636 Reznick, D., Butler, M. J., IV & Rodd, H. (2001) Life-history evolution in guppies VII. 637 The comparative ecology of high and low predation environments. Am Nat 157: 126– 638 140. (doi:10.1086/318627) 639 Sansom, A., Lind, J. & Cresswell, W. (2009) Individual behavior and survival: the roles 640 of predator avoidance, foraging success, and vigilance. Behav Ecol 20, 1168–1174. 641 (doi:10.1093/beheco/arp110)

Schmidt, P. S., Matzkin, L., Ippolito, M. & Eanes, W. F. (2005) Geographic variation in

644

645

646

647

648

649

650

651

652

653

654

655

656

657

658

659

660

661

662

diapause incidence, life-history traits, and climatic adaptation in *Drosophila melanogaster*. Evolution 59: 1721–1732. Schmitz, O. J. (2008) Effects of predator hunting mode on grassland ecosystem function. Science 319: 952–954. (doi:10.1126/science.1152355) Schoener, T. W. (1971) Theory of feeding strategies. Annu Rev Ecol Syst 2: 369–404. Stephan, W. & Li, H. (2006) The recent demographic and adaptive history of *Drosophila* melanogaster. Heredity 98: 65–68. (doi:10.1038/sj.hdy.6800901) Stoks, R., McPeek, M. A. & Mitchell, J. L. (2003) Evolution of prey behavior in response to changes in predation regime: damselflies in fish and dragonfly lakes. Evolution 57: 574-585. Turelli, M. & Hoffmann, A. A. (1991) Rapid spread of an inherited incompatibility factor in California *Drosophila*. Nature 353: 440–442. Wilfert, L. & Jiggins, F. M. (2014) Flies on the move: an inherited virus mirrors *Drosophila melanogaster*'s elusive ecology and demography. Mol Ecol 23: 2093–2104. (doi:10.1111/mec.12709) Worthen, W. B. (1989) Predator-mediated coexistence in laboratory communities of mycophagous *Drosophila* (Diptera: Drosophilidae). Ecol Entomol 14: 117–126. (doi:10.1098/rspb.1998.0559) Yurkovic, A., Wang, O., Basu, A. C. & Kravitz, E. A. (2006) Learning and memory associated with aggression in *Drosophila melanogaster*. Proc Natl Acad Sci U.S.A. 103:

17519-17524. (doi:10.1073/pnas.0608211103)

## Supplemental material

## Table of contents

Supplemental material type	Page numbers
Supporting information	1-2
Supplemental figures	2-7
Supplemental tables	9-14
Videos	14

## **Supporting information**

# Differential response to spiders versus mantids

Because spider and mantid population densities vary by season, we had to temporally segregate the spider assays from the mantid assays. We conducted all spider observations between October and December 2012 and all the mantid observations between March and May 2013. Comparing time allocation and frequencies of occurrences in the predator absent state between the two predator treatments suggest that behavioural modifications were predator induced, and not due to seasonal effects (Figure S5 and S6). Although the assays were carried out under highly controlled conditions, to confirm that predator species-specific behavioural differences were not confounded with seasonal differences in behaviour, we performed 6 additional assays (alternating between spider and mantid treatments) within the span of one week. The control experiments show no evidence of confounding effects of season with *D. melanogaster*'s anti-predator behavioural repertoire (Table S9, S11 and S12 below). Ethograms are shown in Supplement a. Furthermore, to confirm that the disturbance we caused (to the assay

chamber) during the addition of a predator did not confound behavioural responses to the predator, we did 3 "no predator" control assays. For these "no predator" controls, instead of adding a predator to the arena, we caused a mild disturbance (~ to intensity of disturbance caused while adding the predator) without actually adding any predator. We found that disturbance caused during predator addition was not responsible for observed behavioural modifications (Table S10 and S13). Finally, "no predator" controls also ruled our temporal differences in fruit fly activity levels (Table S10 and S13)

# **Supplemental figures**

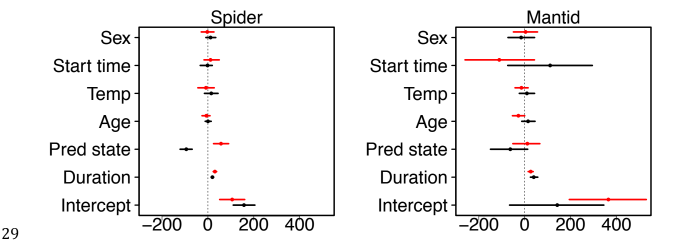
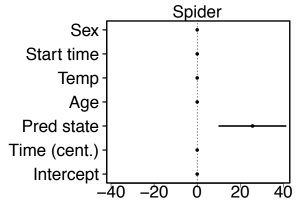


Figure S1

Presence of jumping spiders causes fruit flies to walk more and groom less, whereas the presence of mantids causes weaker, more variable (and not significant) changes in fruit fly activity levels. Here we show coefficient plots from the output of mixed effects models using the package MCMCglmm to visualize duration of two behaviours (**grooming in black** and **locomotion in red**) as a function of predator state (present vs absent of spiders, left panels and mantids, right panels), time spent in the assay [Time (cent.)] total, sex of the fly, start time of the assay, temperature in the room and age of the fly. The continuous covariate "Time" was centered

around the mean and therefore reflects the average increase in time spent per minute of the assay.

Estimates are in seconds. Error bars are  $\pm$  95% CI.



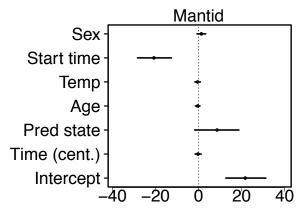


Figure S2

Fruit flies "stop" significantly longer in the presence of spiders (left panel), and to a much lesser extent (and not significantly) in the presence of mantids (right panel). Here we have coefficient plots made from the output of mixed effects models using the package MCMCglmm to visualize duration of "stopping" as a function of predator state (present vs absent), time spent in the assay [Time (cent.)], sex of the fly, start time of the assay, temperature in the room and age of the fly. Estimates are in seconds. Error bars are  $\pm$  95% CI. The continuous covariate "Time" was centered around the mean and therefore reflects the average increase in time spent per minute of the assay. Although assays were performed between 9 am and 12 pm each day, start time for the mantid assays significantly affected the total time that flies spent "stopping".

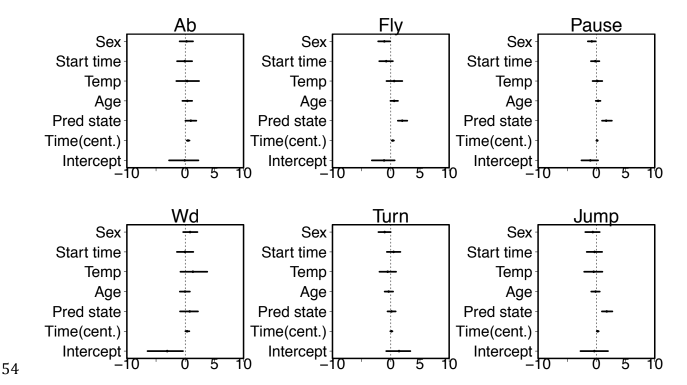


Figure S3

In the presence of spiders, fruit flies increased the frequency with which they performed flights, pauses and jumps. Here we show coefficient plots made from the output of mixed effects models where the events were modeled using a poisson generalized linear mixed model with a log-link function fit using the MCMCg1mm function, and estimates remain on a natural log scale. Coefficient plots were used to visualize frequency of each individual behavioural event (ab, fly, pause, wd, turn and jump) as a function of predator state (present vs absent of a spider), time spent in the assay [Time (cent.)], sex of the fly, start time of the assay, temperature in the room and age of the fly. All estimates are scaled to number of events per minute. Error bars are  $\pm$  95% CI.

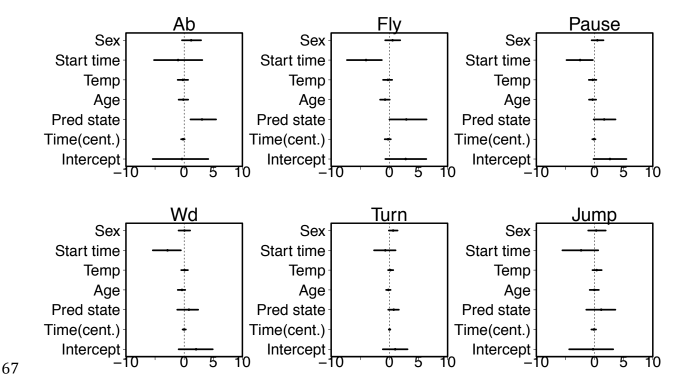


Figure S4

Fruit flies performed abdominal lifts are a higher frequency in the presence of a juvenile mantid. Coefficient plots made from the output of mixed effects models where the events were modeled using a poisson generalized linear mixed model with a log-link function fit using the MCMCglmm function, and estimates remain on a natural log scale. Coefficient of the model output are used to visualize frequency of each individual behavioural event (ab, fly, pause, wd, turn and jump) as a function of predator state (present vs absent of a mantid), time spent in the assay [Time (cent.)], sex of the fly, start time of the assay, temperature in the room and age of the fly. All estimates are scaled to number of events per minute. Error bars are ± 95% CI. Although assays were performed between 9 am and 12 pm each day, start time for the mantid assays significantly affected the frequency at which *D. melanogaster* performed the "Fly", "Wd" and "Jump" behaviours.

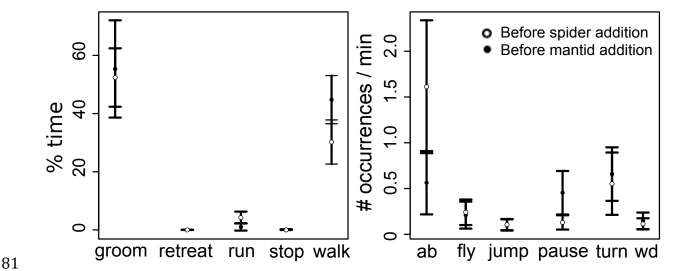


Figure S5

Hunting mode induced behavioural differences in fruit fly behaviours were not confounded with seasonal effects. Here we show percentage time spent in each behavioural state (left) and number of occurrences per minute for each behavioural event (right) as measured for individual fruit flies before the addition of a spider (white circles) and before the introduction of a mantid (black circles) into the chamber. Error bars are  $\pm 2$  \* SEs. Overlapping error bars suggest that there was minimal effect of season on the behavioural repertoire of fruit flies.

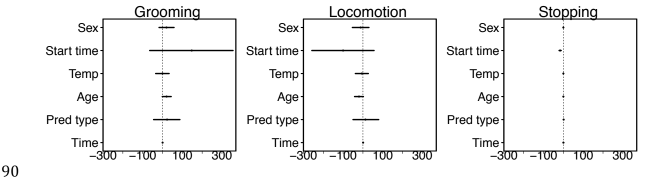
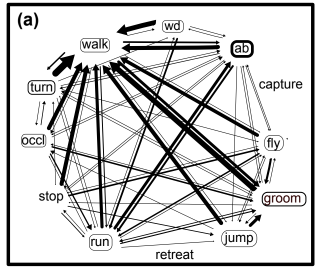


Figure S6

Seasonal differences in fruit fly behaviours did not confound behavioural differences induced by different hunting-modes. Flies measured before the addition of a spider did not differ in behaviour from flies measured before the addition of a mantid.



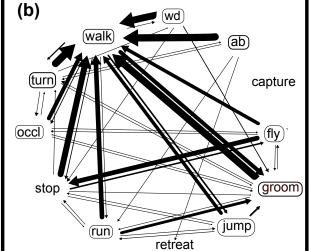


Figure S7

a) A diagram representing probability of transitioning from one fly behaviour to the other when individuals were measured before the addition of a spider b) A diagram representing probability of transitioning from one fly behaviour to the other for individuals measured before the addition of a juvenile mantid. Thickness of arrows indicates transition probability between the two behaviours. The arrowhead points to the behaviour being transitioned to. Thickness of the box around behavioural state ( groom, run, occl, retreat, stop and walk) indicate the mean proportion of total time spent in that behaviour, whereas thickness of the box around behavioural events (fly, jump, turn, wd, ab) indicates mean number of occurrences per minute of that behaviour. To reduce the complexity of the web we combined the behaviours "pause" with the behaviour "stop". Behavioural transitions that occurred less than 10 times have not been shown in the figure.

# **Supplemental tables**

**Table S1** Transition frequency matrix when a spider was present in the chamber. Each row represents the number of times one behaviour (row name) transitioned to another behaviour (column name). Numbers in blue represent transitions that occurred more often that expected under a model of independence, whereas numbers in red are transitions that occurred less often than expected (see methods).

Spider

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd	Total
ab	0	2	8	9	7	0	45	6	5	10	112	7	211
capture	0	0	0	0	0	0	0	0	0	0	0	0	0
fly	4	2	0	15	7	0	34	25	10	1	58	1	157
groom	11	3	16	0	10	0	8	11	5	7	77	2	150
jump	4	2	19	8	0	0	9	13	2	0	28	0	85
retreat	1	0	0	0	0	0	0	0	0	0	0	0	1
run	43	4	33	13	9	1	0	18	3	4	51	9	188
stop	9	3	32	10	16	0	19	0	1	9	38	0	137
occl	3	0	6	4	5	0	4	4	0	1	14	0	41
turn	8	1	0	4	3	0	14	2	0	0	66	2	100
walk	122	7	36	87	26	0	42	60	14	63	0	24	481
wd	5	0	4	0	0	0	9	0	0	1	29	0	48
Total	210	24	154	150	83	1	184	139	40	96	473	45	1599

**Table S2** Transition frequency matrix before a spider was added to the chamber. Each row represents the number of times one behaviour (row name) transitioned to another behaviour (column name). Numbers in blue represent transitions that occurred more often that expected under a model of independence, whereas numbers in red are transitions that occurred less often than expected (see methods)s.

#### No spider

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd	Total
ab	0	0	2	16	1	0	30	1	2	10	65	2	129
capture	0	0	0	0	0	0	0	0	0	0	0	0	0
fly	1	0	0	9	1	0	6	2	0	0	15	0	34
groom	7	0	11	0	8	0	18	1	5	10	122	0	182
jump	0	0	1	7	0	0	0	0	1	0	9	0	18
retreat	0	0	0	0	0	0	0	0	0	0	0	0	0
run	38	0	5	14	1	0	0	4	2	1	42	3	110
stop	1	0	4	0	2	0	2	0	0	1	10	0	20
occl	2	0	0	6	0	0	5	0	0	3	16	0	32
turn	4	0	1	5	0	0	2	0	3	0	72	1	88
walk	74	0	8	126	5	0	42	13	18	64	0	12	362
wd	1	0	0	0	0	0	4	0	0	0	13	0	18
Total	128	0	32	183	18	0	109	21	31	89	364	18	993

**Table S3** Transition frequency in the presence of a juvenile praying mantid. Each row represents the number of times one behaviour (row name) transitioned to another behaviour (column name). Numbers in blue represent transitions that occurred more often that expected whereas numbers in red are transitions that occurred less often than expected.

Mantid

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd	Total
ab	0	1	9	2	1	9	10	5	4	6	127	0	174
capture	0	0	0	0	0	0	0	0	0	0	0	0	0
fly	0	0	0	7	0	1	5	60	2	0	50	0	125
groom	0	0	17	0	3	0	18	3	5	13	305	1	365
jump	0	0	0	1	0	0	0	13	0	0	18	1	33
retreat	9	0	2	0	0	0	0	1	1	7	3	0	23
run	11	0	3	18	0	0	0	17	1	12	68	0	130
stop	4	0	45	9	4	5	21	0	0	5	139	0	232
occl	1	0	3	7	1	0	5	1	0	0	73	0	91
turn	4	1	2	6	1	3	15	0	1	0	164	0	197
walk	145	2	43	326	23	5	51	132	73	150	0	49	999
wd	0	0	1	0	0	0	2	0	0	1	47	0	51
Total	174	4	125	376	33	23	127	232	87	194	994	51	2420

**Table S4** Transition frequency matrix before a juvenile mantid was added to the chamber. Each row represents the number of times one behaviour (row name) transitioned to another behaviour

(column name). Numbers in blue represent transitions that occurred more often that expected whereas numbers in red are transitions that occurred less often than expected.

## No mantid

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd	Total
ab	0	0	0	0	0	1	1	0	0	1	31	0	34
capture	0	0	0	0	0	0	0	0	0	0	0	0	0
fly	0	0	0	1	0	0	0	19	1	0	11	0	32
groom	0	0	9	0	1	0	4	1	3	12	212	0	242
jump	0	0	0	3	0	0	1	4	0	0	6	0	14
retreat	1	0	0	0	0	0	0	1	1	0	0	0	3
run	0	0	0	9	1	0	0	1	0	0	15	0	26
stop	0	0	14	1	1	0	1	0	0	1	54	0	72
occl	0	0	1	2	0	0	0	0	0	1	55	0	59
turn	2	0	0	2	0	0	0	0	4	0	79	0	87
walk	31	0	8	236	10	1	19	45	48	71	0	21	490
wd	0	0	0	2	0	0	0	1	0	0	18	0	21
Total	34	0	32	256	13	2	26	72	57	86	481	21	1080

**Table S5** Transition probability from one behaviour (row name) to the other (column name) in the presence of a zebra jumping spider. Transition probabilities are obtained by dividing each transition frequeny (see table S1) between a pair of behaviours by the total number of times a given behaviour was performed (row sums in table S1).

Spider

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd
ab	0.00	0.01	0.04	0.04	0.03	0.00	0.22	0.03	0.02	0.05	0.55	0.03
capture	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
fly	0.03	0.01	0.00	0.10	0.04	0.00	0.22	0.16	0.06	0.01	0.37	0.01
groom	0.07	0.02	0.11	0.00	0.07	0.00	0.05	0.07	0.03	0.05	0.52	0.01
jump	0.05	0.02	0.22	0.09	0.00	0.00	0.11	0.15	0.02	0.00	0.33	0.00
retreat	1.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
run	0.24	0.02	0.18	0.07	0.05	0.01	0.00	0.10	0.02	0.02	0.28	0.05
stop	0.07	0.02	0.23	0.07	0.12	0.00	0.14	0.00	0.01	0.07	0.28	0.00
occl	0.07	0.00	0.15	0.10	0.12	0.00	0.10	0.10	0.00	0.02	0.34	0.00
turn	0.08	0.01	0.00	0.04	0.03	0.00	0.14	0.02	0.00	0.00	0.67	0.02
walk	0.27	0.02	0.08	0.19	0.06	0.00	0.09	0.13	0.03	0.14	0.00	0.05
wd	0.10	0.00	0.08	0.00	0.00	0.00	0.19	0.00	0.00	0.02	0.60	0.00

**Table S6** Transition probability from one behaviour (row name) to the other (column name) before a zebra jumping spider was introduced into the arena. Transition probabilites are obtained by dividing each transition frequeny (see table S1) between a pair of behaviours by the total number of times a given behaviour was performed (row sums in table S1).

## No spider

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd
ab	0.00	0.00	0.02	0.13	0.01	0.00	0.24	0.01	0.02	0.08	0.51	0.02
capture	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
fly	0.03	0.00	0.00	0.26	0.03	0.00	0.18	0.06	0.00	0.00	0.44	0.00
groom	0.04	0.00	0.06	0.00	0.04	0.00	0.10	0.01	0.03	0.05	0.67	0.00
jump	0.00	0.00	0.06	0.39	0.00	0.00	0.00	0.00	0.06	0.00	0.50	0.00
retreat	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
run	0.36	0.00	0.05	0.13	0.01	0.00	0.00	0.04	0.02	0.01	0.39	0.03
stop	0.05	0.00	0.20	0.00	0.10	0.00	0.10	0.00	0.00	0.05	0.50	0.00
occl	0.06	0.00	0.00	0.19	0.00	0.00	0.16	0.00	0.00	0.09	0.50	0.00
turn	0.05	0.00	0.01	0.06	0.00	0.00	0.02	0.00	0.03	0.00	0.83	0.01
walk	0.21	0.00	0.02	0.36	0.01	0.00	0.12	0.04	0.05	0.18	0.00	0.03
wd	0.06	0.00	0.00	0.00	0.00	0.00	0.22	0.00	0.00	0.00	0.72	0.00

**Table S7** Transition probability from one behaviour (row name) to the other (column name) in the presence of a juvenile praying mantid. Transition probabilites are obtained by dividing each transition frequeny (see table S1) between a pair of behaviours by the total number of times a given behaviour was performed (row sums in table S1).

#### Mantid

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd
ab	0.00	0.01	0.05	0.01	0.01	0.05	0.06	0.03	0.02	0.03	0.73	0.00
capture	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
fly	0.00	0.00	0.00	0.06	0.00	0.01	0.04	0.48	0.02	0.00	0.40	0.00
groom	0.00	0.00	0.05	0.00	0.01	0.00	0.05	0.01	0.01	0.04	0.84	0.00
jump	0.00	0.00	0.00	0.03	0.00	0.00	0.00	0.41	0.00	0.00	0.56	0.03
retreat	0.39	0.00	0.09	0.00	0.00	0.00	0.00	0.04	0.04	0.30	0.13	0.00
run	0.08	0.00	0.02	0.14	0.00	0.00	0.00	0.13	0.01	0.09	0.52	0.00
stop	0.02	0.00	0.19	0.04	0.02	0.02	0.09	0.00	0.00	0.02	0.60	0.00
occl	0.01	0.00	0.03	0.08	0.01	0.00	0.05	0.01	0.00	0.00	0.80	0.00
turn	0.02	0.01	0.01	0.03	0.01	0.02	0.08	0.00	0.01	0.00	0.83	0.00
walk	0.15	0.00	0.05	0.34	0.02	0.01	0.05	0.14	0.08	0.16	0.00	0.05
wd	0.00	0.00	0.02	0.00	0.00	0.00	0.04	0.00	0.00	0.02	0.92	0.00

**Table S8** Transition probability from one behaviour (row name) to the other (column name) before a juvenile praying mantid was introduced into the arena. Transition probabilites are obtained by dividing each transition frequeny (see table S1) between a pair of behaviours by the total number of times a given behaviour was performed (row sums in table S1).

#### No mantid

Behavior	ab	capture	fly	groom	jump	retreat	run	stop	occl	turn	walk	wd
ab	0.00	0.00	0.00	0.00	0.00	0.03	0.03	0.00	0.00	0.03	0.91	0.00
capture	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
fly	0.00	0.00	0.00	0.03	0.00	0.00	0.00	0.59	0.03	0.00	0.34	0.00
groom	0.00	0.00	0.04	0.00	0.00	0.00	0.02	0.00	0.01	0.05	0.88	0.00
jump	0.00	0.00	0.00	0.21	0.00	0.00	0.07	0.29	0.00	0.00	0.43	0.00
retreat	0.33	0.00	0.00	0.00	0.00	0.00	0.00	0.33	0.33	0.00	0.00	0.00
run	0.00	0.00	0.00	0.35	0.04	0.00	0.00	0.04	0.00	0.00	0.58	0.00
stop	0.00	0.00	0.19	0.01	0.01	0.00	0.01	0.00	0.00	0.01	0.75	0.00
occl	0.00	0.00	0.02	0.03	0.00	0.00	0.00	0.00	0.00	0.02	0.93	0.00
turn	0.02	0.00	0.00	0.02	0.00	0.00	0.00	0.00	0.05	0.00	0.91	0.00
walk	0.07	0.00	0.02	0.50	0.02	0.00	0.04	0.10	0.10	0.15	0.00	0.04
wd	0.00	0.00	0.00	0.10	0.00	0.00	0.00	0.05	0.00	0.00	0.86	0.00

**Table S9** Proportion of time spent in a given behavioural state by each individual fruit fly before and after introducing a treatment (i.e., a disturbance, spider or mantid) to the assay chamber.

		Groo	ming	Walking		Running		Stopping		Retreat	
Treatment	Individual	Absent	Present	Absent	Present	Absent	Present	Absent	Present	Absent	Present
	1	0.820	0.728	0.180	0.265	0.000	0.005	0.000	0.000	0.000	0.000
Disturbance	2	0.590	0.265	0.338	0.658	0.000	0.009	0.044	0.015	0.000	0.000
	3	0.264	0.219	0.658	0.685	0.048	0.036	0.013	0.052	0.000	0.000
	1	0.528	0.490	0.359	0.394	0.022	0.011	0.000	0.003	0.000	0.000
Spider	2	0.116	0.000	0.844	0.861	0.000	0.086	0.000	0.053	0.000	0.000
	3	0.706	0.181	0.262	0.368	0.010	0.076	0.000	0.347	0.005	0.002
	1	0.419	0.764	0.474	0.217	0.008	0.005	0.061	0.014	0.000	0.000
Mantid	2	0.673	0.075	0.327	0.785	0.000	0.014	0.000	0.093	0.000	0.006
	3	0.684	0.963	0.309	0.037	0.000	0.000	0.000	0.000	0.000	0.000

**Table S10** Number of occurrences per minute of each behavioural event before and after the introduction of a treatment (i.e., a disturbance, spider or mantid) to the assay chamber.

		Pause		Turn		Ab		Wd		Fly		Jump	
Treatment	Individual	Absent	Present										
	1	0.000	0.006	0.000	0.006	0.000	0.003	0.000	0.000	0.000	0.002	0.000	0.003
Disturbance	2	0.003	0.013	0.003	0.013	0.060	0.023	0.003	0.005	0.082	0.027	0.000	0.005
	3	0.000	0.002	0.000	0.002	0.027	0.052	0.003	0.010	0.045	0.090	0.018	0.016
	1	0.009	0.013	0.000	0.005	0.043	0.046	0.006	0.000	0.006	0.003	0.000	0.000
Spider	2	0.020	0.018	0.032	0.045	0.012	0.000	0.012	0.027	0.048	0.252	0.000	0.000
	3	0.015	0.011	0.003	0.011	0.026	0.090	0.000	0.002	0.018	0.041	0.003	0.000
	1	0.052	0.017	0.016	0.006	0.031	0.002	0.008	0.002	0.070	0.020	0.003	0.000
Mantid	2	0.005	0.060	0.005	0.012	0.023	0.111	0.000	0.003	0.000	0.076	0.000	0.006
	3	0.000	0.000	0.003	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000

169 Table S11 Coefficients from a linear model (lm) for control individuals meansured before and after170 the addition of a spider. While estimate of posterior means are similar to those of the main spider171 dataset, Due to low sample sizes, CIs are large.

	Grooming	1	
Coefficient	Posterior mean	Lower CI	Upper Cl
Intercept	181.88	-23.34	387.10
Predator state	-75.38	-377.82	227.05
Duration	0.45	-0.34	1.24
	Locomotio	n	
Intercept	156.76	37.32	276.21
Predator state	17.27	-158.75	193.29
Duration	0.24	-0.22	0.70

**Table S12** Coefficients from a linear model (lm) for control individuals meansured before and 175 after the addition of a mantid. Estimate of posterior means are similar to those of the main mantid 176 dataset, but due to low sample sizes, CIs large.

Grooming				
Coefficient	Posterior mean	Lower CI	Upper Cl	
Intercept	134.79	-1399.38	1668.96	
Predator state	341.56	-2661.10	3344.23	
Duration	-0.80	-12.76	11.17	
Locomotion				
Intercept	345.69	-919.99	1611.36	
Predator state	-323.02	-2800.19	2154.15	
Duration	1.64	-8.24	11.51	

179 **Table S13** Coefficients from a linear model (lm) for control individuals meansured before and after 180 a disturbance. Despite low sample sizes, it is clear that disturbance had minimal effect on fruit fly 181 behaviours.

Grooming				
Coefficient	Posterior mean	Lower CI	Upper CI	
Intercept	200.00	60.11	339.81	
Disturbance	0.00	-197.78	197.78	
Duration	2.38	-0.53	5.29	
Locomotion				
Intercept	132.30	-0.28	264.84	
Disturbance	0.00	-187.47	187.47	
Duration	-1.05	-3.81	1.70	

# 184 Links to videos describing novel behaviours

178

- 185 1) Abdominal Lifting http://dx.doi.org/10.6084/m9.figshare.1185638
- 186 2) Stopping Behaviour http://dx.doi.org/10.6084/m9.figshare.1185639
- 187 3) Retreat http://dx.doi.org/10.6084/m9.figshare.1185640