1	TITLE: Mechanisms of reduced interspecific interference between territorial species
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17	ABSTRACT: Interspecific territoriality has complex ecological and evolutionary
18	consequences. Species that interact aggressively often exhibit spatial or temporal shifts in
19	activity that reduce the frequency of costly encounters. We analyzed data collected over a
20	13-year period on 50 populations of rubyspot damselflies (Hetaerina spp.) to examine
21	how rates of interspecific fighting covary with fine-scale habitat partitioning and to test
22	for agonistic character displacement in microhabitat preferences. In most sympatric

23 species, interspecific fights occur less frequently than expected based on the species' 24 relative densities. Incorporating measurements of spatial segregation and species 25 discrimination into the calculation of expected frequencies accounted for most of the 26 reduction in interspecific fighting (subtle differences in microhabitat preferences could 27 account for the rest). In 23 of 25 sympatric population pairs, we found multivariate 28 differences between species in territory microhabitat (perch height, stream width, current 29 speed, and canopy cover). As predicted by the agonistic character displacement 30 hypothesis, sympatric species that respond more aggressively to each other in direct 31 encounters differ more in microhabitat use and have higher levels of spatial segregation. 32 Previous work established that species with the lowest levels of interspecific fighting 33 have diverged in territory signals and competitor recognition through agonistic character 34 displacement. In the other species pairs, interspecific aggression appears to be maintained 35 as an adaptive response to reproductive interference, but interspecific fighting is still 36 costly. We now have robust evidence that evolved shifts in microhabitat preferences also 37 reduce the frequency of interspecific fighting.

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39 KEYWORDS: agonistic character displacement, habitat partitioning, habitat preference,
40 microhabitat, interference competition, interspecific territoriality, Odonata

INTRODUCTION

43	Competition between animal taxa is widespread and often involves aggression.
44	Interspecific aggression may initially arise from misdirected intraspecific aggression
45	(Murray 1971; Nishikawa 1987; Singer 1989; Schultz and Switzer 2001; Korner et al.
46	2010; Cowen et al. 2020). However, in the absence of a contested resource, species are
47	expected to diverge in ways that reduce the frequency and costs of interspecific
48	aggression, a process known as divergent agonistic character displacement (ACD). Most
49	documented cases of divergent ACD involve evolutionary shifts in agonistic signals and
50	competitor recognition (Grether et al. 2009; Grether et al. 2013; Grether et al. 2017;
51	Latour and Ganem 2017; Moran and Fuller 2018a; Moran and Fuller 2018b; Zambre et
52	al. 2020). Conversely, species with a contested resource in common may converge in
53	agonistic signals and competitor recognition to facilitate resource defense and
54	partitioning, i.e., convergent ACD (Cody 1973; Grether et al. 2009; Reif et al. 2015;
55	Pasch et al. 2017; Souriau et al. 2018; Kirschel et al. 2019). Interspecific mate
56	competition arising from reproductive interference has also been shown to cause
57	convergent ACD (Drury, Okamoto, et al. 2015; Grether et al. 2020).
58	In addition to convergence or divergence in traits related to competitor
59	recognition, interspecific aggression can also cause drastic changes in species' spatial and
60	temporal niches (Melville 2002; Gotelli et al. 2010; Pigot and Tobias 2013; Edgehouse et
61	al. 2014; Ulrich et al. 2017; Eurich et al. 2018). One common result of interspecific
62	territoriality is competitive displacement where a dominant or more aggressive species
63	forces a subordinate species into a different habitat or to be active during different

64	periods (Garcia 1983, Reitz and Trumble 2002, Langkilde and Shine 2004, Jankowski et
65	al. 2010, Pasch et al. 2013, Kajtoch et al. 2015). In other cases, both species may shift in
66	habitat use or temporal activity (Bay et al. 2001; Eurich et al. 2018; Reif et al. 2018).
67	Whether the shift in activity occurs in one or both species, interspecific competition is
68	reduced, although one species may be forced into suboptimal habitat (Randall 1978;
69	Robinson and Terborgh 1995; Grether et al. 2013). Habitat partitioning can occur at
70	various spatial and temporal scales, such as elevational or latitudinal gradients on a
71	macroscale (Connell 1983; Schoener 1983; Lewin 1989; Hawkins 1999; Mark et al.
72	2001) as well as across small scale variation in microhabitat (Eurich et al. 2018; Reif et
73	al. 2018).

74 Habitat partitioning could also arise from species differences in habitat 75 preferences that evolved in response to selection against interspecific fighting in the past, 76 which would be a form of divergent ACD (Grether et al. 2009). As yet, however, there 77 are few if any well documented cases of ACD in habitat preferences (for a possible 78 example, see Vallin et al. 2012). Species can diverge in habitat use for many reasons, and 79 determining whether these differences are products of past or ongoing interspecific 80 interactions is challenging (Connell 1978; Ross 1986; Wisheu 1998; Pinter-Wollman et 81 al. 2006).

Rubyspot damselflies (*Hetaerina*) are a good system for examining the
relationship between interspecific aggression and niche partitioning because levels of
interspecific aggression vary widely among sympatric species. Male rubyspots defend
mating territories along streams and rivers (Johnson 1963; Córdoba-Aguilar et al. 2009;

86	Anderson and Grether 2011; but see Guillermo-Ferreira and Del-Claro 2011). Mature
87	males of all species in the genus have red pigmentation at the base of their wings,
88	whereas the lack of red or other conspicuous pigmentation in female wings makes them
89	more cryptic (Garrison 1990). Simulated territory intrusion and wing color manipulation
90	experiments have shown that the responses of territory holders to intruders is largely
91	based on wing coloration and that species with more similar wing coloration respond
92	more aggressively to each other (Anderson and Grether 2010a; Drury and Grether 2014;
93	Grether et al. 2015).
94	Interspecific aggression in <i>Hetaerina</i> appears to largely be an adaptive response
95	to interspecific mate competition (Drury, Okamoto, et al. 2015; Drury, Anderson, et al.
96	2019; Grether et al. 2020), but selection should still favor adaptations that reduce the
97	frequency of interspecific fighting. Territorial fights are costly, primarily because they
98	can result in males losing their territories and priority of access to ovipositing females
99	(territory possession confers a three-fold mating advantage; Grether 1996; Drury &
100	Grether 2014). Damselfly fights also have energetic and physiological costs (reviewed in
101	Suhonen et al. 2008; Vieira and Peixoto 2013; Córdoba-Aguilar and González-Tokman
102	2014; Kemp 2018; Grether 2019), and fights that do not immediately result in territory
103	turnover likely reduce the ability of the residents to win future fights. Thus, selection may
104	favor divergence in microhabitat use because this reduces the probability of interspecific
105	encounters and therefore the frequency of interspecific fights.
106	Species differences in microhabitat use have been documented in Hetaerina

107 (Johnson 1973; Anderson and Grether 2011), but it is unknown whether these differences

are products of past or ongoing interspecific aggression. Sympatric species could differ in
microhabitat use for reasons unrelated to interspecific aggression, or because one species
actively displaces the other from preferred microhabitat, or because of selection against
interspecific fighting, i.e., divergent ACD (Grether et al. 2009).

112 In this paper, we analyze data collected in the field over a 13-year period on 14 113 species pairs of rubyspot damselflies to examine whether current levels of interspecific 114 fighting can be explained by species pair differences in spatial segregation and species 115 discrimination, and to test the ACD prediction that species that respond more 116 aggressively to each other in direct encounters show higher levels of divergence in 117 microhabitat use. To the best of our knowledge, this is the first study, on any taxon, to 118 approach either of these questions with an adequate level of replication at the species 119 level.

120

121 **METHODS**

122 Sympatric populations

123 We studied 14 sympatric species pairs (10 different species) across 15 sites in the

124 southwestern United States, Mexico, and Costa Rica from 2005 to 2017 (see

125 Supplementary Methods S1 for criteria for inclusion of study sites). Some sites were

126 visited multiple times in different years. Because of interannual variation in microhabitat

- 127 availability and species densities, pooling the data across visits could have obscured
- 128 patterns of interest. We therefore kept visits to the same sites in different years separate

for the analyses presented in this paper, for a total of 25 species-pair-site-years, which werefer to henceforth as sympatric population pairs (Table S1).

131

132 **Behavioral sampling**

At the beginning of each study period, we established a 200-300 m transect by fastening a 133 134 rope with numbered flags in 1-m increments along one or both banks of the river. Males 135 within the transect were captured with aerial nets, marked on the abdomen with unique 136 color codes using paint pens (200-S Fine Point, MarvyDecocolor Paint Marker; Uchida 137 of America, Torrance, CA, USA; Anderson et al. 2011), photographed (Figure S1), and 138 released where they were captured. Each day, 2-5 observers continuously walked along 139 the transect during periods of territorial activity (~0800-1800) recording the ID and 140 locations of males (as [x, y, z] coordinates where x is the flag number, y is the distance 141 from the bank of the stream, and z is the height) to the nearest 0.1 m on hand-held 142 computers (Psion PLC, London). In total, we marked 7,483 males and made 34,614 143 observations. A male was considered a territory holder if he was observed in the same 144 location $(\pm 2 \text{ m})$ and perching close to the water, as males do when they are defending a 145 site, on at least two consecutive days (Anderson and Grether 2010a). 146 When fights were observed, we recorded the species involved, the males' IDs (if 147 marked), and the location. Fights between the same two marked males on the same day, 148 and fights involving unmarked males at the same location on the same day, were

149 collapsed into a single fight for the purpose of calculating intra- and interspecific fighting

150 frequencies (Anderson and Grether 2011; Drury and Grether 2014).

151

152 Microhabitat sampling

153 To quantify the microhabitat use of the damselflies at each site, we measured canopy 154 cover, stream width, stream current speed, and the perch height of territory holders. We 155 measured canopy cover, an especially important axis of microhabitat variation for 156 ectothermic insects (Shelly 1982; Huey 1991; Tsubaki et al. 2010; Okuyama et al. 2013), 157 every 5-10 m along the stream using a concave spherical densiometer (Forestry 158 Suppliers, Inc.). We made one measurement in the middle of the stream where the stream 159 was <3 m wide, two measurements (one on each side of the stream) where the stream was 160 3-10 m wide, and three measurements (one on each side of the stream and another in the 161 middle) where the stream was ≥ 10 m wide. Canopy cover ranges from 0 to 100% and 162 higher values indicate shadier habitat. We measured stream width every 2 m along the 163 transect with a measuring tape and visually assessed current speed every 2 m near both 164 banks and the middle of the stream using a 0 to 4 scale, where 0 is still water and 4 is 165 rapidly moving white water. To characterize the microhabitat of each male's territory, we 166 interpolated between the two nearest canopy cover, stream width, and current speed 167 readings, and averaged the male's recorded perch heights.

168

169 **Expected frequencies of interspecific fighting**

170 We considered interspecific fighting to be reduced relative to intraspecific fighting if the 171 observed frequency of interspecific fights was lower than expected based on a simple null

172 model (Anderson and Grether 2011). The null model assumes that males encounter and 173 fight with conspecific and heterospecific males in direct proportion to the species' 174 relative densities. The null expectation for the frequency of interspecific fights is simply 175 the observed total number of fights multiplied by $2d_id_j$ where d_i and d_j are the species' 176 relative densities (Supplementary Methods S2a). All previous comparisons of observed 177 and expected rates of interspecific fighting in *Hetaerina* were based on this null model 178 (Anderson and Grether 2011; Drury et al. 2015).

179 Factors that could cause the observed frequency of interspecific fights to be lower 180 than the null expectation include: (1) spatial segregation between the species, which 181 would reduce the frequency of interspecific encounters; and (2) species discrimination, which would reduce the probability of interspecific encounters resulting in territorial 182 183 fights. To evaluate whether spatial segregation alone accounts for the reduction in 184 interspecific fighting, we constructed lists of all males of each species observed within 4 185 m of the center of each established territory. From these lists, we calculated the average 186 proportion of heterospecific "neighbors" from each species' perspective and multiplied 187 the average of these two estimates by the total number of observed fights to obtain the 188 expected frequency of interspecific fights (Supplementary Methods S2b). The 4 m-189 criterion is based on the observation that males respond to conspecific males up to $\sim 2 \text{ m}$ 190 away from their perch and in doing so could enter the reaction zone of a male perched 4 191 m away; beyond a distance of 4 m males are unlikely to interact (Anderson and Grether 192 2011 used the same criterion for similar reasons).

193	To evaluate whether species discrimination alone accounts for the reduction in
194	interspecific fighting, we calculated the expected interspecific fighting rate by
195	multiplying the null expectation by the probability of males chasing a heterospecific
196	intruder. Heterospecific chase probabilities were calculated from the results of simulated
197	territory intrusion tests in which territory holders were presented with live, tethered males
198	in timed trials (Anderson and Grether 2010a; Supplementary Methods S2c).
199	Finally, we combined the methods above to calculate the expected frequency of
200	interspecific fighting based on observed levels of spatial segregation and species
201	discrimination (Supplementary Methods S2d).
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203	Heterospecific aggression ratio
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212 We used chi-squared tests to determine whether observed frequencies of interspecific

- 213 fighting differed from expected frequencies, the Monte Carlo simulation method to
- calculate *p*-values in cases with expected frequencies < 5, and Holm's sequential

215 Bonferroni procedure (Holm 1979) to correct for multiple comparisons. Wilcoxon

216 matched pairs signed ranks tests were used to compare the alternative methods of

217 calculating expected frequencies to the null model.

To examine the causes of variation in interspecific fighting rates, we constructed a linear multiple regression model where the dependent variable was the log of the ratio of the observed number of interspecific fights to the expected number of fights under the null model. The predictors were the mean proportion of heterospecific neighbors, the species difference in microhabitat use, and the heterospecific chase probability (n = 25;

223 Supplementary Methods S2c Equation S2b).

We used principal component analysis (PCA) to find the principal axes of variation in microhabitat use across all territory holders in the study (n = 1974). To obtain an overall measure of the species difference in microhabitat use at each site, we calculated the Euclidean distance between the species' PC centroids (n = 25).

228 To more fully characterize species differences in microhabitat use at each site, we 229 used linear discriminant analysis (LDA) and mixed-effects multivariate regression. The 230 microhabitat variables were transformed to a mean of 0 and variance of 1 to make the 231 LDA coefficients comparable and to weight the microhabitat variables equally in the 232 regression models. The predictor variables in the regression models were species (1 or 2), 233 an index identifying the microhabitat variable (1-4), the species by microhabitat variable 234 interaction, and a random-effects term for male ID (n = 1974). To make the sign of the 235 mean difference between species the same for all four microhabitat variables, we

assigned the species with the smaller mean an index of 1 and the species with the largermean an index of 2 (Table S3).

238 The ACD hypothesis predicts that species that respond more aggressively when 239 interspecific encounters occur should differ more in microhabitat use and exhibit higher 240 levels of spatial segregation. To test the first part of this prediction, we constructed linear 241 mixed-effects regression models with all sympatric population pairs included (n = 25)242 and nested random-effects terms for population pair and male ID. The full, multivariate 243 model included indices to identify the species (1 or 2) and microhabitat variables (1-4), 244 the mean HA ratio, and all interactions between these terms. We also constructed 245 separate models for each microhabitat variable, with species index, mean HA ratio, the 2-246 way interaction, and a random-effects term for sympatric population pair, and used the 247 Akaike information criterion (AIC) to evaluate whether adding quadratic terms improved 248 the model fit. We used a similar approach to test for effects of heterospecific aggression 249 on the proportion of heterospecifics in a male's territory neighborhood but coded the 250 species index based on the relative density of territory holders (1 = low, 2 = high). 251 Mixed-effects regression models were implemented with mixed in STATA 16.1 252 (StataCorp 2019). Other analyses were carried out in R 4.0.3 – 4.0.5 (R Core Team 2020, 253 2021); LDA was implemented with the lda default in R package MASS 7.3-53.1 254 (Venables and Ripley 2002). 255

256 **RESULTS**

257 Species differences in microhabitat use

258 The first three principal components (PCs) accounted for 83.4% of the variance in 259 microhabitat use (Table 1). PC1 explained 33.8% of the variance and had a large positive 260 loading for canopy cover and negative loadings for the other variables (Table 1; Figure 261 S2). PC2 explained 26.3% of the variance and had a large negative loading for current 262 speed and a positive loading for perch height, while PC3 explained 23.3% of the variance 263 and had a large positive loading for stream width and a negative loading for current speed 264 (Table 1; Figure S2). PC1 likely represents variation in stream size (smaller streams tend 265 to be slower and make smaller gaps in the forest canopy) while the other axes represent 266 variation in stream gradient and size independent of canopy cover (males tend to perch 267 low on emergent rocks in fast current and higher in the bank vegetation in slower 268 sections).

Twenty three of the 25 sympatric populations differ significantly in microhabitat use (Table 2). Overall, the LDA correctly classified 79.7% of territory holders to species based on microhabitat use, and for many populations the species classifications were 80-100% correct (Table 2). As shown by the species means and LDA coefficients, all four microhabitat variables proved useful for differentiating between sympatric species (Tables 2, S3).

275

276 Interspecific fighting

Across the 25 pairs of sympatric populations, we collected data on 1,974 territory holders and 1,793 fights, of which 346 (19.3%) were between heterospecific males. The observed frequency of interspecific fights was significantly lower than the null expectation in 21

280 out of 25 cases (Table 3). There was considerable variation in this relationship across 281 species, as reflected by the wide range of chi-square values (Table 3). The multiple 282 regression analysis with species differences in neighborhood composition, microhabitat, 283 and chase probabilities as predictors accounted for 54% of the variation in the ratio of 284 observed to expected interspecific fights (Table 4). The greater the proportion of 285 heterospecifics in a territory holder's neighborhood and the greater the species difference 286 in microhabitat use, the lower the ratio of observed to expected interspecific fights. 287 We were also able to explain much of the reduction in the frequency of observed 288 interspecific fights compared to the null expectation. In some sympatric population pairs, 289 the reduction in the frequency of interspecific fights was explained by spatial segregation 290 (Figure 1a), while in others the reduction was explained by species discrimination (Figure 291 1b) or both spatial segregation and species discrimination (Figure 1c). Yet there were 292 some sympatric populations for which these factors could not fully explain the reduction 293 in observed interspecific fights (Figure 1d). The mean difference between the number of 294 observed and expected interspecific fights decreased by 32.4% when the expected rate 295 was calculated using only neighborhood composition, 19.1% using only chase 296 probabilities, and 50% with neighborhood composition and chase probabilities combined 297 (Table 5).

298

299 Effects of interspecific aggression on microhabitat and spatial partitioning

300 Overall, we found striking support for the hypothesis that interspecific aggression drives

301 species apart in microhabitat use. In the full multivariate model, the 3-way interaction

302	was highly significant ($\chi^2 = 85.70$, df = 3, $p < 0.0001$), which indicates that the effect of
303	heterospecific aggression on the species difference in microhabitat use varies strongly
304	among microhabitat variables. We therefore analyzed the microhabitat variables
305	separately. Adding quadratic terms substantially improved the fit of the perch height
306	($\Delta AIC = -15.55$) and stream width ($\Delta AIC = -14.58$) models but worsened the fit of the
307	current speed ($\Delta AIC = 3.25$) and canopy cover models ($\Delta AIC = 0.26$). The species
308	difference in perch height was greater between sympatric populations with low and high
309	HA compared to those with intermediate HA (Figure 2; species XHA: -0.68 \pm 0.14, z = -
310	4.91, $p < 0.0001$; species×HA ² : 0.43 ± 0.11, $z = 3.96$, $p = 0.001$). The species difference
311	in the other three microhabitat variables increased with the HA ratio (Figure 2; stream
312	width, speciesXHA: -4.64 \pm 1.88, z = -2.47, p = 0.013; speciesXHA ² : 5.95 \pm 1.45, z =
313	4.11, $p < 0.0001$; current speed, species×HA: 0.16 ± 0.074, $z = 2.20$, $p = 0.028$; canopy
314	cover, speciesXHA: 15.31 \pm 2.28, $z = 6.70$, $p < 0.0001$). Also as predicted by the ACD
315	hypothesis, the proportion of heterospecific neighbors decreased, and thus spatial
316	segregation increased, with the level of heterospecific aggression (Figure 3; HA: -0.20 \pm
317	0.06, $z = -3.22$, $p = 0.001$), particularly for species with a low relative density of territory
318	holders, as indicated by a positive interaction between the relative density of territory
319	holders and the HA ratio (0.095 \pm 0.035, $z = 2.72$, $p = 0.0066$).
320	To evaluate whether the results were affected by males at site GO contributing
321	data to two different sympatric population pairs, we ran the mixed-effects regression
322	models on subsets of the data and found that dropping any two GO pairs had no
323	qualitative effect on the results (Table S4).

345

Grether et al. 2022).

325 **DISCUSSION**

326 This 13-year investigation of 14 species pairs provides an unprecedented level of support 327 for the general hypothesis that interspecific aggression increases spatial habitat 328 partitioning between sympatric species. Specifically, we found that sympatric species that 329 are more aggressive to each other in simulated intruder tests differ more in microhabitat 330 use (Figure 2) and are more spatially segregated (Figure 3). In principle, three non-331 mutually exclusive mechanisms could have produced this pattern: species sorting, 332 competitive displacement, and agonistic character displacement (ACD). We discuss each 333 of these potential mechanisms in turn and explain why we consider ACD to be the most 334 likely mechanism. 335 In this context, species sorting refers to effects of interspecific interactions on the 336 probability of species occurring in sympatry (Pfennig and Pfennig 2012). If interspecific 337 fighting reduces the probability of co-occurrence, the positive relationship between 338 microhabitat partitioning and heterospecific aggression could be a byproduct of variation 339 in the level of microhabitat divergence prior to secondary contact. It has yet to be shown, 340 however, that interspecific fighting affects the probability of co-occurrence in 341 damselflies. Most research on coexistence mechanisms in Odonata has focused on 342 resource competition and predation at the larval stage (e.g., McPeek 2004; Siepielski et 343 al. 2010; Siepielski et al. 2011; Bried and Siepielski 2019); it is not yet clear whether 344 behavioral interference at the adult stage affects coexistence in this taxon (reviewed in

346 Competitive displacement has been shown, or strongly inferred, to be the primary 347 cause of species differences in habitat use in other territorial animals, including insects 348 and arachnids (Reitz and Trumble 2002), barnacles (Connell 1961), mammals (Brown 349 1971; Pasch et al. 2013), birds (Garcia 1983; Jankowski et al. 2010; Kajtoch et al. 2015; 350 Martin and Bonier 2018), and reptiles (Langkilde and Shine 2004; Edgehouse et al. 351 2014). A common feature of systems in which competitive displacement occurs is that 352 one species is competitively superior and displaces the other species from the preferred 353 habitat (Reitz and Trumble 2002). In general, there are several ways that one species 354 could be competitively superior, but in the case of damselflies competing for mating 355 territories, competitive superiority would entail behavioral dominance or superior aerial 356 fighting ability. We are not aware of any rubyspot damselfly species pairs in which one 357 species is dominant or consistently wins territorial fights, but further research is 358 warranted. Whether competitive displacement occurs, and the extent to which it explains 359 the effects of heterospecific aggression on microhabitat use, could be tested with removal 360 experiments or microhabitat manipulations.

While species sorting and competitive displacement are both plausible post-hoc explanations, neither of those hypotheses could have been used to predict that microhabitat partitioning would correlate positively with heterospecific aggression without making unsupported assumptions about the study system. By contrast this was a well-founded prediction of the agonistic character displacement hypothesis. Previous research showed that some sympatric rubyspot damselfly species have diverged substantially in male wing coloration and competitor recognition, and that the territories

368	of these species often overlap extensively (Anderson and Grether 2010a; Anderson and
369	Grether 2010b; Anderson and Grether 2011). In most rubyspot damselfly species pairs,
370	however, interspecific territorial aggression is adaptive because females of these species
371	are too similar in coloration for males to distinguish between them; a territory holder that
372	tolerated heterospecific males on his territory would risk losing mating opportunities
373	(Drury, Okamoto, et al. 2015; Drury et al. 2019; Grether et al. 2020). In this situation,
374	divergence in microhabitat preferences might be the only way for selection to reduce the
375	costs of interspecific aggression. Our initial evidence that microhabitat divergence has
376	evolved in response to interspecific aggression was based on four sympatric species pairs
377	(Anderson and Grether 2011). Now with data on 14 sympatric species pairs, across
378	multiple sites and years, we can confirm that microhabitat divergence is strongly
379	associated with interspecific aggression (Figure 2).
380	We expect positive relationships between habitat partitioning and heterospecific
381	aggression to be found in other taxa as well. Our other findings are rather damselfly
382	specific, but parallels might be found in other taxa. For example, interspecifically
383	territorial birds are expected to vertically stratify in habitats with a large height
384	dimension, such as woodlands (Murray 1971). Indeed lunulated and Salvin's antbirds
385	(Gymnopithys lunulate and G. salvini) forage from taller perches in the presence of
386	larger, behaviorally dominant antbirds and woodcreepers (Willis 1968). Similarly, the
387	iguanid lizard Liolaemus tenuis perches higher when sympatric with the aggressively
388	dominant L. pictus (Medel et al. 1988). Rubyspot damselfly species with both low and
389	high levels of heterospecific aggression differ more in mean perch height than those with

intermediate levels of heterospecific aggression (Figure 2). Considering that species with
low levels of heterospecific aggression have overlapping territories (Anderson & Grether
2011), the species differences in perch height probably function to reduce accidental
interspecific interference.

394 We found a negative relationship, at the population level, between the mean 395 proportion of heterospecific neighbors and the ratio of observed to expected frequencies 396 of interspecific fights (Table 4). Logically, territory holders with more heterospecific 397 neighbors should be observed in more, not fewer, interspecific fights. The 398 counterintuitive population-level result is probably an artifact of the mathematical 399 constraint that males in populations with lower relative densities have more 400 heterospecific neighbors. What this population-level analysis did show, however, is that 401 species differences in microhabitat use reduce interspecific fighting (Table 4). 402 When we based expected frequencies on the proportion of heterospecific 403 neighbors and the probability of males responding aggressively to heterospecifics, the 404 mean difference between the observed and expected numbers of interspecific fights was 405 50% less than under the null model (Table 5). However, the observed number of 406 interspecific fights was still significantly below the expected number in many populations 407 (Figures 1, S3). Species differences in microhabitat use, which were found in all but two 408 sympatric population pairs (Table 2), likely reduce the frequency of interspecific fights 409 below what would be expected based on the composition of territorial neighborhoods and 410 heterospecific aggression. For example, species that perch at different heights tend to 411 fight at different heights (authors, pers. obs.), and therefore may be less likely to fight

412 with heterospecific neighbors than expected based on the spatial arrangement of 413 territories. Additionally, differences between species in stream current speed preferences 414 might also reduce the frequency of interspecific fighting because current speed can vary 415 among neighboring territories. 416 Studies on other taxa have also revealed adaptive connections between 417 interspecific aggression and microhabitat use. For example, fine-scale microhabitat 418 partitioning has been reported in interspecifically territorial damselfish (Eurich et al. 419 2018). Territorial neotropical cichlid fish (*Amphilophilous* spp.) are more likely to 420 tolerate heterospecific neighbors with divergent coloration (Lehtonen et al. 2010; 421 Lehtonen et al. 2015). Interspecifically aggressive nightingales (Luscinia megarhynchos 422 and L. luscinia) "escape" to allotopic sites in the sympatric region of their geographic 423 ranges and occupy habitat avoided by the congener (Reif et al. 2018). 424 Selection against interspecific interference is only one of many possible reasons 425 that closely related species might differ in microhabitat preferences. For example, 426 microhabitat preferences could have diverged in allopatry before secondary contact 427 (Berner and Thibert-Plante 2015; Dufour et al. 2015). Conspecific attraction might also 428 reduce spatial overlap between sympatric species (Scott and Lee 2013; Stodola and Ward 429 2017) and result in chance differences in microhabitat use (Buxton et al. 2020). 430 Nevertheless, the results presented here provide compelling evidence that interspecific 431 aggression has played an important role in microhabitat divergence. 432

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451 DATA ACCESSIBILITY

452 Analyses reported in this article can be reproduced using the data provided by McEachin et453 al. (2021).

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674	

676	FIGURE LEGENDS
677	Figure 1. Observed and expected numbers of interspecific fights, based on different
678	methods of calculating the expectation, for selected sympatric population pairs (see Fig.
679	S3 for the remaining populations). An example of the reduction in the number of
680	observed interspecific fights that can be explained by (a) spatial segregation of territory
681	holders based on the proportion of heterospecific males in territory holders'
682	neighborhoods (H. occisa-H. miniata at LH 2016), (b) competitor recognition based on
683	the aggressive response of territory holders to heterospecific intruders relative to
684	conspecific intruders in simulated territory intrusions (H. occisa-H. titia at OT 2007), (c)
685	both spatial segregation and competitor recognition (H. fuscoguttata-H. titia at GO 2016),
686	and (d) a population pair in which neither spatial distribution nor competitor recognition
687	can account for the reduction in observed interspecific fights (H. cruentata-H. majuscula
688	at SL 2016). Combined refers to the model that calculates expected fighting rates based
689	on both spatial segregation and competitor recognition. See Table 3 for statistical results
690	comparing all 25 sympatric population pairs.
691	

692 **Figure 2.** Evidence that interspecific aggression causes species to diverge in

693 microhabitat preferences. As heterospecific aggression increases, so do species

694 differences in territory microhabitat. The exception is perch height, which differs the least

695 between sympatric species at intermediate levels of heterospecific aggression. Points and

bars represent population means and standard errors. Triangles (circles) represent the

697	population with higher (lower) relative density in each pair. Colors uniquely identify the
698	paired populations. Lower (upper) black lines represent predicted values for the
699	populations with lower (higher) means of the corresponding microhabitat variable, and
700	gray areas are 95% confidence intervals, from the mixed-effects regression model
701	described in Statistical methods .
702	
703	Figure 3. Evidence that interspecific aggression increases spatial separation between
704	species. The proportion of heterospecific neighbors decreases as heterospecific
705	aggression increases. The slope of the relationship is steeper for populations with low
706	relative density compared to those with high relative density. Lower (upper) black lines
707	represent predicted values for the populations with lower (higher) relative density in each
708	pair. All other symbols and codes follow Figure 2.

	PC1 (33.8%)	PC2 (26.3%)	PC3 (23.3%)	PC4 (16.6%)
Perch height	-0.579	0.522	-0.014	-0.626
Stream width	-0.345	-0.380	0.858	-0.017
Current speed	-0.284	-0.760	-0.459	-0.361
Canopy cover	0.682	-0.066	0.231	-0.691

Table 1 Microhabitat principal component loadings (% variance explained)

Table 2 Results of linear discriminant analyses (LDA) and multivariate regression 714

	LDA species classif.					LDA coefficients (standardized)			
Pop.	Spp.	#	#	%	Perch	Stream	Current	Canopy	
pair	code	correct	wrong	correct	height	width	speed	cover	Z.
1	c	41	9	82	-0.31	-0.07	-1.40	0.50	7.62***
	0	33	5	87					
2	0	45	10	82	0.77	0.10	-0.30	0.23	4.83***
	t	17	12	59					
3	0	11	27	29	0.66	-0.02	-0.94	-0.30	2.23*
	t	44	10	81					
4	а	15	14	52	0.79	-0.20	-1.10	0.70	5.70***
	t	68	3	96					
5	а	38	9	81	0.78	-0.19	-1.17	0.66	4.82***
	t	18	16	53					
6	а	14	4	78	0.98	-0.03	0.06	-0.18	2.64**
	t	24	3	89					
7	а	48	8	86	-0.10	-0.01	0.65	1.06	5.85***
	с	23	4	85					
8	c	54	0	100	0.75	0.64	0.87	2.48	1.20
	v	0	15	0					
9	а	92	5	95	1.36	-0.04	-0.11	0.87	4.88***
	v	8	19	30					
10	0	120	19	86	2.17	-0.15	-0.09	0.32	5.61***
	t	38	38	50					
11	0	190	12	94	2.36	-0.11	-0.18	0.39	6.37***
	t	27	31	47					
12	0	46	3	94	1.73	-0.03	0.23	2.82	4.07***
	S	12	10	55					
13	0	16	3	84	-0.14	-0.95	0.17	6.15	2.22*
	р	9	3	75					
14	n	6	0	100	-1.03	0.20	-0.16	-0.85	14.65***
	0	80	2	98					
15	с	10	18	36	1.13	0.59	-0.76	-1.16	3.21**
	m	64	4	94					

analyses (MVA) of species differences in microhabitat use 715

16	c	28	8	78	0.83	0.35	0.53	2.14	6.32***
	р	34	0	100					
17	0	29	18	62	2.29	0.01	0.10	4.98	2.45*
	р	42	10	81					
18	n	8	8	50	2.32	0.03	-0.45	-1.48	2.35*
	t	48	2	96					
19	n	19	3	86	-0.28	0.11	-0.03	-0.61	6.34***
	0	41	12	77					
20	f	6	1	86	-2.25	0.39	1.79	-0.83	3.11*
	0	10	0	100					
21	f	4	3	57	0.51	-0.20	-0.50	-0.85	3.81***
	t	19	1	95					
22	0	8	2	80	0.80	-0.42	-1.02	-0.29	3.98***
	t	20	0	100					
23	f	16	2	89	-1.50	0.06	0.30	-2.19	2.38*
	0	11	6	65					
24	f	18	0	100	0.45	-0.24	-0.03	-2.48	1.27
	t	4	6	40					
25	0	13	4	76	1.87	-0.23	-0.49	0.66	2.35*
	t	6	4	60					

716 See Table S1 for site info and species names.

717 *p < 0.05, **p < 0.01, ***p < 0.001

	Observed freq. Expected freq.					
Pop.		Intrasp.	Intersp.	Intrasp.	Intersp.	Chi-sq.
pair	Spp.	fights	fights	fights	fights	test
1	c	8	2	2.8	9.0	15.78**
	0	9		7.3		
2	0	13	6	19.4	25.3	88.18***
	t	34		8.3		
3	0	8	4	8.6	26.3	42.00***
	t	43		20.1		
4	a	6	8	2.4	18.6	12.63*
	t	43		36.0		
5	а	25	18	19.1	30.1	9.44*
	t	18		11.9		
6	а	3	3	3.5	18.7	23.86**
	t	41		24.8		
7	а	37	7	31.0	19.0	17.91*
	c	9		2.9		
8	c	38	23	39.9	23.5	4.46
	v	7		3.4		
9	а	31	12	28.5	16.2	2.57
	V	4		2.3		
10	0	16	11	15.6	21.2	17.47**
	t	17		7.2		
11	0	30	14	19.2	41.0	35.55***
	t	38		21.8		
12	0	26	12	17.5	26.5	15.61***
	S	16		10.0		
13	0	25	5	15.4	19.5	20.57***
	р	11		6.2		
14	n	2	2	2.2	25.5	29.25**
	0	98		74.3		
15	c	29	26	12.6	60.2	45.36***
	m	90		72.2		
16	с	27	15	4.5	33.8	123.95**

719720 Table 3 Comparison of the observed frequencies of interspecific fights to those expected

121 under the num model

	р	60		63.7		
17	0	42	24	32.8	69.9	69.05***
	р	74		37.2		
18	n	7	33	13.2	75.8	49.12***
	t	158		109.0		
19	n	4	6	4.2	19.2	17.36**
	0	35		21.6		
20	f	13	7	10.2	10.2	1.85
	0	3		2.6		
21	f	13	18	10.1	36.7	17.81**
	t	49		33.2		
22	0	3	4	15.4	27.9	134.86***
	t	49		12.7		
23	f	145	53	125.5	100.4	64.22***
	0	48		20.1		
24	f	145	18	132.7	35.0	18.94*
	t	7		2.3		
25	0	48	15	39.6	26.1	8.20
	t	7		4.3		

722 See Table S1 for site info.

723 *p < 0.05, **p < 0.01, ***p < 0.001 after sequential Bonferroni correction

Table 4 Predictors of variation in the ratio of observed to expected interspecific fights

Predictor	В	SE	р
Intercept	0.01	0.24	0.98
Neighborhood composition	-1.08	0.39	0.011
Microhabitat differences	-0.30	0.058	< 0.001
Chase probabilities	0.29	0.15	0.078

value the null model

728 Linear multiple regression, n = 25, model adjusted $R^2 = 0.54$, F(3, 21) = 10.4, p < 0.001.

730731 Table 5 Comparison of three alternative methods of calculating expected frequencies of

732	interspecific fighting to the null model, with Wilcoxon matched pairs signed ranks tests
	Obs. – Exp.

	Obs. –	Obs. – Exp.		
Model	Mean	SE	V	р
Relative density (null)	-18.8	2.52	-	-
Spatial segregation	-12.7	2.33	57	0.0034
Species discrimination	-15.2	2.37	57	0.0065
Spatial segregation and species	-9.4	2.04	30	< 0.001
discrimination combined				

n = 25 population pairs











