1	Male sex pheromone components in the butterfly <i>Heliconius</i>				
2	melpomene				
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10	Abstract				
11	Sex specific pheromones are known to play an important role in butterfly courtship, and may				
12	influence both individual reproductive success and reproductive isolation between species.				
13	Extensive ecological, behavioural and genetic studies of Heliconius butterflies have made a				
14	substantial contribution to our understanding of speciation. However, although long suspected				
15	to play an important role, chemical signals have received relatively little attention in this				
16	genus. Here, we combine morphological, chemical and behavioural analyses of a male				
17	pheromone in the neotropical butterfly Heliconius melpomene. First we identify specialized				
18	brush-like scales that are putative androconia, and lie within the shiny grey region found on				
19	the hindwing of males. We then describe six putative male sex pheromone compounds, which				
20	are largely confined to the androconial region of the hindwing of mature males, but not				
21	immature males or females. Finally, behavioural assays reveal subtle, but detectable,				
22	differences in female response to models scented with hindwing androconial extracts of				
23	mature conspecific males as compared to unscented controls. Collectively, the results describe				
24	structures involved in release of the pheromone and a list of potential male sex pheromone				
25	compounds triggering a behavioural response in females.				
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27	Key words: Heliconius, pheromone, androconia, Lepidoptera, sexual selection.				
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30 Introduction

Organisms use chemicals for a wide variety of forms of communication. In particular, sex 31 32 pheromones are species-specific blends of chemical compounds mediating intraspecific communication between males and females, and can play key roles in both determining the 33 34 reproductive success of individuals within a species, and in reproductive isolation between species (Wvatt, 2003, 2014). Some insects deploy single chemicals as signals, but in many 35 36 insects, pheromone communication is dependent on complex combinations of chemical components (Grillet, Dartevelle, & Ferveur, 2006; Nieberding et al., 2008; Symonds, 37 Johnson, & Elgar, 2012). This chemical complexity provides the potential to convey 38 sophisticated information (Nieberding et al., 2012). The best studied insect sex pheromones 39 are perhaps the chemicals produced by female moths to attract mating partners, often over 40 long distances (Löfstedt, 1993; Smadja & Butlin, 2008). Male insects can also produce sex 41 pheromones but these have generally received less attention (Wyatt, 2014). 42 Whilst variation in sex pheromone blend is now known to be a major determinant of 43 reproductive isolation and speciation in many species of moths (Löfstedt, 1993; Smadja & 44 Butlin, 2008; Lassance et al., 2010), to date sex pheromones have been studied in just a few 45 species of butterfly. Acceptance behaviour in the queen butterfly Danaus berenice is 46 47 regulated by a dihydropyrrolizine alkaloid released by the male (Brower & Jones, 1965; Meinwald, Meinwald, & Mazzocchi, 1969; Pliske & Eisner, 1969). Another danaine butterfly, 48 49 Idea leuconoe displays 'hair-pencils' during courtship, which contain a mixture of dihydropyrrolizine alkaloids, aromatics, terpenoids, hydrocarbons and a series of γ -lactones 50 51 (Nishida *et al.*, 1996). This volatile mixture applied on dummy male butterflies elicits an acceptance posture in females. Pieris rapae and Pieris brassicae use macrocyclic lactones as 52 a pheromone to induce acceptance in females (Yildizhan et al., 2009). In Bicvclus anvnana 53 males with reduced amounts of male sex pheromone have decreased mating success implying 54 55 a direct involvement in reproductive fitness (Nieberding et al., 2008, 2012). Male wing 56 compounds are also known to contribute to reproductive isolation between closely related species of butterflies (Grula, McChesney, & Taylor, 1980; Phelan & Baker, 1987; Bacquet et 57 al., 2015). 58 Here we focus on the potential role of a male contributed pheromone in *Heliconius* 59 butterflies. Heliconius is a diverse neotropical genus, studies of which have contributed 60 greatly to our understanding of speciation (Jiggins, 2008; Merrill et al., 2015). These 61 butterflies are known for Müllerian mimicry, where unrelated species converge on the same 62 warning signal to more efficiently advertise their unpalatability to predators. However, 63

64 closely related *Heliconius* taxa often differ greatly in colour pattern and divergent selection

acting on warning patterns is believed to play an important role in speciation within the genus

(Bates, 1862; Jiggins *et al.*, 2001; Merrill *et al.*, 2011b). In particular, males are known to use
colour pattern to recognize mates (Jiggins *et al.*, 2001; Jiggins, Estrada, & Rodrigues, 2004;
Kronforst *et al.*, 2006; Melo *et al.*, 2009; Merrill *et al.*, 2011a; Merrill, Chia, & Nadeau, 2014;

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Finkbeiner, Briscoe, & Reed, 2014).

70 Some aspects of chemical signaling have been studied in *Heliconius*. For example, males are known to transfer an antiaphrodisiac pheromone to females during mating (Gilbert, 71 72 1976). Once mated, females produce a strong odor that acts to repel approaching males (Schulz et al., 2008). Studies in H. melpomene have shown that abdominal glands of males 73 contain a complex odor bouquet consisting of the volatile compound (E)- β -ocimene together 74 with some trace components and esters of common C16 – and C18 – fatty acids with alcohols, 75 where β -ocimene acts as the main antiaphrodisiac pheromone. This bouquet is formed during 76 first few days of eclosion and transferred during copulation to the females (Schulz et al., 77 2008). This antiaphrodisiac effect can be observed in several Heliconius species, which show 78 79 species-specific patterns of scent gland constituents (Gilbert, 1976). In addition, Heliconius may also use green leaf volatiles during mate searching. Six-carbon alcohols and acetates are 80 released in larger amounts after tissue damage by caterpillars, which males of the pupal 81 82 mating species *H. charithonia* then use to find potential mates (Estrada & Gilbert, 2010). 83 Once males find pupae they also use chemical cues to determine sex (Estrada et al., 2010).

There are also a number of observations that indicate a role for chemical recognition 84 85 during adult mating. H. erato males can distinguish between wings dissected from conspecific and heterospecific females that are virtually identical in wing pattern, but this effect 86 87 disappears after wings have been washed in hexane (Estrada & Jiggins, 2008). H. cvdno males show a preference for their own pattern over that of the closely related *H. melpomene*, 88 89 but will court wing pattern models of *H. melpomene*. However, *H. cvdno* males have virtually never been observed mating with *H. melpomene* females, suggesting strong barriers in 90 91 addition to wing pattern (Naisbit, Jiggins, & Mallet, 2001). In addition, H. melpomene coexists with visually almost identical co-mimics, notably H. erato, which is likely to favour 92 the evolution of pheromonal recognition signals to avoid confusion in courtship. 93 In some populations the very closely related species *H. timareta*, is sympatric and 94

mimetic with *H. melpomene*, but nevertheless displays strong assortative mating (Giraldo *et al.*, 2008). A recent study of sympatric *H. melpomene* and *H. timareta* in Peru has provided
some of the first evidence for a role of chemical signals in species recognition (Mérot *et al.*,
2015). Experiments with perfumed males using abdominal scent glands and wing extracts
shows increased probability of inter-specific mating when males were perfumed with
heterospecific extracts. In addition, chemical analysis of both abdominal glands and whole

wings provided evidence for differences between these closely related species in their
chemical signatures (Mérot *et al.*, 2015)

103 Here we focus on the wing pheromones of *H. melpomene*. First, we investigate morphological structures potentially associated with pheromone production. In butterflies, a 104 105 variety of species-specific structures including brushes, fans, and differentiated scales on wings, legs or abdomen are used to expose pheromones produced in associated glands (Wvatt, 106 107 2003; Nieberding et al., 2008). In particular, male specific scent glands, termed androconia 108 are common across the Lepidoptera. In male Heliconius, a patch of shiny grey scales is present on the overlapping region of the hind and forewing (Figure 1). The observed sexual 109 dimorphism in this trait suggests that these are androconia, and may be associated with a male 110 sex pheromone (Emsley, 1963). Furthermore, earlier authors have identified brush-like scales 111 112 in this region that are the putative site for pheromone production (Müller, 1912; Barth, 1952). Here we investigate the structure of these scales using Scanning Electron Microscopy. 113 Second, we complement recently published chemical analysis of whole *H. melpomene* wings 114 by dissecting wing regions to identify those wing regions that are associated with the 115 production of compounds and identify potential male sex pheromone compounds isolated 116 from this region. Finally, using a simple behavioural assay we evaluate female response to 117 118 male androconial pheromone extracts.

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120 Methods

Individuals used for morphological and chemical analyses were from an outbred stock of 121 122 Heliconius melpomene plesseni and Heliconius melpomene malleti (sold as H. m. aglaope) maintained at the University of Cambridge insectaries (Figure 1). These two races are from 123 124 the region of a hybrid zone in the eastern Andes of Ecuador, and showed signs of inter-racial hybridization in the stocks, so are treated here as a single population and referred to as the 125 126 Ecuador samples. These stocks were established from individuals obtained from a commercial 127 breeder (Stratford Butterfly Farm: www.butterflyfarm.co.uk). Laboratory stocks were maintained on the larval food plants, Passiflora menispermifolia and Passiflora biflora. Adult 128 butterflies were fed on ~10% sugar solution mixed with an amino acid supplement (Critical 129 Care Formula[®]). In addition, five individual males of *H. m. rosina* were sampled from the 130 wild around the town of Gamboa in Colón Province, Panama, and are referred to as the 131 Panama samples. 132

133

134 Morphological analysis

135 The detailed morphology of androconial scales was determined using Field Emission

136 Scanning Electron Microscope. Three males and two females of *H. melpomene* from Ecuador

137 were used for this analysis. The androconial grey scale region was dissected out from both

138 hind and forewings and attached to aluminium stubs with carbon tabs and subsequently

139 coated with 20nm of gold using a Quorum/Emitech sputter coater. The gold-coated

140 androconia were then viewed in an FEI XL30 FEGSEM operated at 5kV. Images were

141 recorded digitally using XL30 software at 500x magnification.

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143 Characterization of potential male sex pheromone

144 Wing tissue from ten male (five newly emerged and five 10-day old mature individuals) and five female (10-day old) individuals from the Ecuador stock was collected between November 145 2011 and March 2012 for chemical analysis. In addition, wing tissue from five male adult 146 individuals from Panama was collected in August 2012 for chemical analysis. Wings were 147 dissected into four parts: forewing androconia, hindwing androconia, forewing non-148 androconia and hindwing non-androconia. The 'androconia' regions corresponded to the 149 grey-brown region shown in Figure 1c, with non-androconia corresponding to the remaining 150 portion of the wing. In females, a region corresponding in size and extent to the grey-brown 151 region seen in males was dissected. The dissected sections were then allowed to soak in either 152 hexane or dichloromethane for 3 hours in a glass vial. The solvent was then transferred to new 153 154 vial and stored at -20°C. Initial tests showed no major differences between hexane and 155 dichloromethane extracts. Therefore, the more polar dichloromethane was used in later 156 analysis.

Extracts were analyzed by gas chromatography/mass spectrometry (GC/MS) using a 157 158 Hewlett-Packard model 5975 mass-selective detector connected to a Hewlett-Packard GC model 7890A, and equipped with a Hewlett-Packard ALS 7683B autosampler. A HP-5MS 159 160 fused silica capillary column (Agilent, $30 \text{ m} \times 0.25 \text{ mm}$, $0.25 \text{ }\mu\text{m}$) was used. Injection was performed in splitless mode (250°C injector temperature) with helium as the carrier gas 161 162 (constant flow of 1.2 ml/min). The temperature programme started at 50°C, was held for 5 min, and then rose to 320°C with a heating rate of 5°C/min. All components were identified 163 by comparison of mass spectra and gas chromatographic retention index with those of 164 authentic reference samples, and analysis of mass spectra. The double bond position of 165 unsaturated compounds were determined by derivatisation with dimethyl disulfide (Buser et 166 al., 1983). The alcohols were synthesised from the corresponding esters by reduction 167 according to established procedures (Becker & Beckert, 1993, p. 570). The aldehydes were 168 synthesised by oxidation of the respective alcohols (More & Finney, 2002). Synthesis of the 169 methyl branched alcohols and aldehydes will be reported elsewhere (F. Mann et al., in 170 preparation). The samples were quantified by using gas chromatography with flame ionisation 171 172 detection with a Hewlett-Packard GC model 7890A equipped with a Hewlett-Packard ALS

- 173 7683B autosampler. A BPX-5 fused silica capillary column (SGE, $25 \text{ m} \times 0.22 \text{ mm}$, $0.25 \mu \text{m}$)
- 174 was used. Injection was performed in splitless mode (250°C injector temperature) with
- 175 hydrogen as the carrier gas (constant flow of 1.65 ml/min). The temperature programme
- started at 50°C, held for 5 min, and then rose to 320°C with a heating rate of 5°C/min.
- 177 Pentadecyl acetate (10.1 ng) or (*Z*)-4-tridecenyl acetate (1 ng) were used as internal standard.
- 178 Only compounds eluting earlier than hexacosane were considered for analysis. Later
- 179 compounds were identified as cuticular hydrocarbons, 2,5-dialkyltetrahydrofurans, cholesterol
- 180 and artefacts like phthalates or adipates. The variability in the late eluting cuticular
- 181 hydrocarbons was low and did not show characteristic differences.
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183 Behavioural experiments

184 Behavioural assays to determine female response to the putative male sex pheromone were

- 185 conducted in the *Heliconius* insectaries at the Smithsonian Tropical Research Institute's
- 186 (STRI) in Gamboa, Panama between January 2013 and July 2013. Females were assayed
- 187 from an outbred stock of *H. melpomene rosina* established from wild individuals collected in
- 188 Gamboa (9°7.4′ N, 79°42.2′ W, elevation 60 m) and the nearby Soberanía National Park,
- 189 República de Panamá. Stocks were maintained on the larval food plants, Passiflora
- 190 *menispermifolia* and *Passiflora biflora*. Adult butterflies were fed on ~20% sugar solution
- 191 with pollen supplements and *Psychotria sp.* and *Psiguiria sp.* serving as an additional pollen
- 192 source. Males sacrificed for extracts for behavioural experiments were either from this same
- 193 stock or were wild individuals collected in Gamboa or the Soberanía National Park. Male
- 194 extracts were obtained by dissecting the androconial region from both hindwings, followed by
- soaking in 200 µl of hexane for 2-3 hours before transferring the solvent to a new glass vial.
- 196 This vial was sealed and stored at -20°C until required.

Behavioural experiments were performed in cages approximately 2m x 2m x 2m. 197 198 Ten virgin females (<5 days after eclosion) were presented with a paper model of H. melpomene rosina attached to a ~30cm length of wire, itself attached to a ~60cm stick on 199 which a camera (Xdreme HD 1080P Action camera) was mounted to record female 200 behaviour. Pheromone extracts from individual H. melpomene males or a control (solvent 201 only) were applied to a strip of Whatman filter paper no. 4, which was attached to the paper 202 butterfly model. The extract of a single male was used for each individual female, so in total 203 ten virgin females and ten male extracts were used. Each trial proceeded as follows: After 204 mounting the paper model on the wire, 200 µl of extract/control was applied to the filter 205 paper. Trials were conducted blind: A third party applied the extract/control so that the 206 experimenter did not know whether they were presenting the focal female with the extract or 207 208 control. The paper model was then presented to the focal female, and the apparatus

209 manipulated to simulate the hovering flight observed in courting males. Each presentation 210 lasted one minute, followed by a one-minute break when the model was removed from the 211 vicinity of the focal female. This was repeated five times before a further five-minute pause in 212 the trial. The whole procedure was repeated six times, alternating between the two treatments, 213 so that each female experienced a total of 15 presentations with the male extract and 15 214 presentations with the control (solvent only).

215 Female behaviours were scored from videos recorded during the trials. To avoid 216 observer bias, the names and order of the digital video files were randomized and only 217 reassigned to treatment and individual once scoring was complete. We determined the behaviours to be recorded from observations of interactions between H. melpomene males and 218 219 females (Table 1). These were 'Slow and moderate wing flapping', where females opened and closed their wings slowly at regular intervals displaying their wing colors; 'Flying/Flying 220 facing towards model'where females tried to chase the paper models for short intervals or 221 females hovered facing the model; 'Slow rhythmic flight' where females displayed slow flight 222 with intermittent gliding; 'Wing display with abdomen normal' where females opened their 223 wings displaying the dorsal wing surface with the abdomen between the wings; 'Quick and 224 jerky wing flapping' where there was continuous wing fluttering by females that effectively 225 226 prevented close contact by the model; 'Flying away from model' where females flew away from the paper model avoiding any interaction; 'Fast erratic flight' where such flight was 227 228 notably very fast, abrupt and apparently directionless and finally 'Wings open with abdomen erect' where females opened their wings displaying the dorsal wing surface with the abdomen 229 230 erect. Each of these behaviours was scored as having occurred or not occurred during each of the one-minute presentations. This led to a dataset for individual females where the proportion 231 232 of one-minute presentations in which each behaviour was observed was calculated (for male 233 extract and control trials separately). To reduce the number of dependent variables we then 234 performed a principal component analysis on these data. The principal component scores for 235 each individual were extracted and were used to test for differences in response between presentations with the male extract and control (solvent only) in paired tests. Statistical 236 237 analyses were performed with *R* (version 3.1.2).

238

239 Results

240 Morphological analysis

In order to investigate structural differences in scale morphology potentially associated with pheromone production, we observed androconial scales of *H. melpomene rosina* males and females under the scanning electron microscope (Figure 2). We identified a marked sexual dimorphism in scale structure. In the central region of the male hindwing androconia along 245 vein Sc+R1 we identified scales with brush-like structures (Figure 2d) at their distal end (Figure 2a). These scales were completely absent in females and in the forewing androconia 246 247 of the males (Figure 2). These scales were also not detected in any other wing region 248 examined. The brush-like scales were found in alternating rows with scales with a normal 249 structure. Moving away from the Sc+R1 wing vein, the density and width of these scales decreased, with isolated brush-like scales found completely surrounded by normal scales. In 250 251 addition, it was found that the base of these brush-like scales was more swollen and glandular 252 as compared to other scales, perhaps indicating a location for the storage or production of the 253 pheromone (Figure 3).

254

255 Characterization of potential male sex pheromone

Using gas chromatography linked to mass spectrometry (GC/MS) and synthesis, six potential 256 male sex pheromone components were found in the male wing extracts from Ecuador samples 257 (Figure 4). These compounds were, (Z)-9-octadecenal, octadecanal, henicosane, (Z)-11-258 icosenal, icosanal and (Z)-13-docosenal. Out of these six compounds, only henicosane was 259 found in all regions of the wing, but (Z)-9-octadecenal, octadecanal, (Z)-11-icosenal, icosanal 260 and (Z)-13-docosenal were restricted to the androconial scale region of the wing. The 261 262 hindwing androconial region tended to contain higher titres of these five compounds as compared to forewing androconial wing region (Figure 5). Octadecanal was present in higher 263 264 amounts in hindwing androconia, while icosanal were only found in trace quantities. (Z)-9-Octadecenal was found in small amounts only in some of the Ecuador samples. A related 265 266 compound, (Z)-11-Icosanol, was found in trace amounts in two Ecuador mature males. Except for henicosane, all five compounds were observed to be age-specific and sex-specific. 267 268 Henicosane was present in the hindwing androconia of both 10-day old and newly emerged males and in females of all ages. In contrast, (Z)-9-octadecenal, octadecanal, (Z)-11-icosenal, 269 270 icosanal and (Z)-13-docosenal were only present in extracts from the hindwing and roconia of 271 10-day old males. The chemical analysis of the extracts of the Panama samples showed a similar composition, although they contained more compounds in slightly higher 272 concentrations. Major components of both the Panama and Ecuador individuals were 273 octadecanal, (Z)-11-icosenal, icosanal, and (Z)-13-docosenal. (Z)-11-Icosenol was found in 274 larger amounts in the Panama samples. They additionally contained high amounts of 275 octadecanol. Small amounts of nonadecanal, methyl-branched octadecanals and their 276 respective alcohols occurred as well. Some of them were identified to be 15-, 16-, and 17-277 methyloctadecanals and the respective alcohols. These compounds were found only in the 278 279 Panama samples with the exception of 17-methyloctadecanal, which was present in two of the 280 samples from Ecuador. *n*-Alkanes were not present in these extracts.

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282 Behavioural experiments

- 283 We tested ten individual virgin females in our behavioural trials, each of which was presented
- sequentially with either male wing extract or a control (solvent only) applied to a paper
- 285 model. Behavioural scores were analysed using a principal components analysis, with two
- principal components retained that together accounted for 80.38% of the variance in the data
- 287 (Table 1, see also supplementary file 1 for the raw data). The second axis accounted for
- 18.3% of the variance. There was no significant difference between male extract and control
- (solvent only) trials for PC1 (Figure 6; Wilcoxon signed rank test: V = 33, p > 0.05);
- 290 however, there was a significant difference between male extract and control (solvent only)
- trials for PC2 (Figure 6; Wilcoxon signed rank test: V = 49, p < 0.05).
- 292

293 Discussion

Many previous studies of *Heliconius* butterflies have described a role for visual cues in mate finding, courtship behaviour, reproductive isolation and speciation (Crane, 1955; Brown, 1981; Jiggins *et al.*, 2001; Kronforst *et al.*, 2006; Giraldo *et al.*, 2008; Estrada & Jiggins, 2008; Merrill *et al.*, 2011a). Here, we have identified compounds associated with sexually mature male wings and described morphological structures putatively involved in pheromone release. Furthermore, we have shown that wing extracts influence female behaviour.

300 We identified six potential MSP compounds from male wing extracts, namely, (Z)-9octadecenal, octadecanal, henicosane, (Z)-11-icosenal, icosanal and (Z)-13-docosenal. Five of 301 302 these are restricted to the androconial region of hindwings of 10-day old males, suggesting a likely role in courtship. Our results are broadly comparable with another recent analysis of wing 303 304 compounds in Heliconius (Mérot et al., 2015), although this recent study did not compare 305 different wing regions, or similarly aged males and females. As the study also did not use 306 synthesis to identify compounds, our work is highly complementary and extends their results to confirm localization of compounds to older males. Male Heliconius do not become sexually 307 active until several days after eclosion, so the absence of these compounds from females and 308 309 younger males is strongly suggestive of a role in mating behaviours. That these five MSP compounds are largely restricted to the hindwing androconia of mature males (Figure 5a) 310 suggests that pheromone storage or production is restricted to the hindwing. Trace amounts on 311 the forewing androconia may be due to contact in the overlapping portion of the fore- and 312 hindwings, and both wings may play a role in dispersal of the compounds during courtship. The 313 wild samples from Panama had both a greater diversity of compounds and higher concentrations 314 on their wings. Further work will be needed to determine whether this is characteristic of natural 315

populations, or reflects a difference between geographic populations of *H. melpomene*, or is
perhaps simply a result of natural inter-individual variation.

318 Scanning electron microscope images of the androconia support storage and/or production of MSPs in the hindwing. The androconial region of male hindwings are equipped 319 320 with special brush-like scales (Figure 2a), which might facilitate the release of the pheromone during courtship and are completely absent in females and any other region of male wings. 321 322 These were located primarily around and along the hindwing vein Sc+R1, similar to the 323 depiction in Figure 73 of Emsley's previous morphological analysis (Emsley, 1963). Similar scales have been described from light microscopy in other Heliconius species, but not 324 previously in *H. melpomene* (Müller, 1912; Barth, 1952). The base of these special brush-like 325 scales was more swollen and glandular as compared to other scales (Figure 3), perhaps 326 indicating a role in storage or production of pheromones by these scales. Even though 327 Heliconius do not have the dramatic sexual dimorphism in hair pencils or brush-like androconia 328 seen in other butterflies, they nonetheless do possess male-specific structures likely associated 329 with pheromone production. 330

We were able to detect a significant, though subtle, difference in the behavioural 331 332 response of females towards models treated with male wing extracts as compared to an 333 unscented control. This indicates that females both detect compounds found on the male wings, and alter their behaviour in response to those compounds. This supports our hypothesis that 334 335 these compounds act as a pheromone involved in courtship behaviour. Nonetheless, it remains unclear exactly what the information is that is conveyed by these signals. The signal may 336 337 influence female courtship, although it is likely that these compounds convey complex information about male species identity, quality, age etc. that are interpreted by females in 338 combination with visual and tactile cues. Recent experiments carried out in Peru also 339 340 demonstrate the role of chemical signals in species recognition between *H. melpomene* and *H.* 341 *timareta*, although these do not distinguish between the role of wing and abdominal pheromone signals (Mérot et al., 2015). Further experiments are necessary to disentangle the details of 342 inter-specific signaling during courtship in Heliconius. We are also currently unable to 343 determine which of the compounds identified here are biologically active. Experiments with 344 synthetic blends of putative MSP compounds were largely inconclusive, so are not presented 345 here. 346

The use of multiple signals is common in animal communication (Candolin, 2003). Moths being nocturnal mainly depend upon chemical cues to attract their mates (Ando, Inomata, & Yamamoto, 2004). On the other hand, butterflies primarily use visual cues to locate mates (Kemp & Rutowski, 2011). It has been shown in *B. anynana* that in addition to visual cues, chemical cues also play a role and are equally important in sexual selection by female

352 choice (Costanzo & Monteiro, 2007). Divergent color patterns and extensive mimicry by

- 353 different species of *Heliconius* has resulted in the use of multiple signals to maintain species
- 354 specificity. Two sympatric cryptic species that share a wing pattern, *H. melpomene malleti* and
- 355 *H. timareta florencia*, nonetheless show strong assortative mating. This suggests that enhanced
- 356 divergence in pheromonal signals acts as an important cue in reproductive isolation in these
- 357 species (Giraldo *et al.*, 2008). Exploring the pheromonal signals and morphological structures
- involved in the process in *H. melpomene* is our first step towards understanding multimodal
- 359 signaling and its role in reproductive isolation.
- 360

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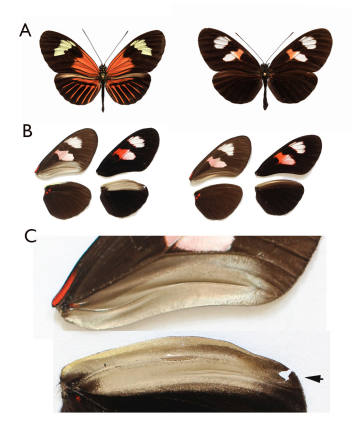
494 Figure 1. A) *H. melpomene malleti* male (left) and *H. melpomene plesseni* female (right). B)

495 Dissected wings from specimens of *H. melpomene plesseni* showing sexual dimorphism in the

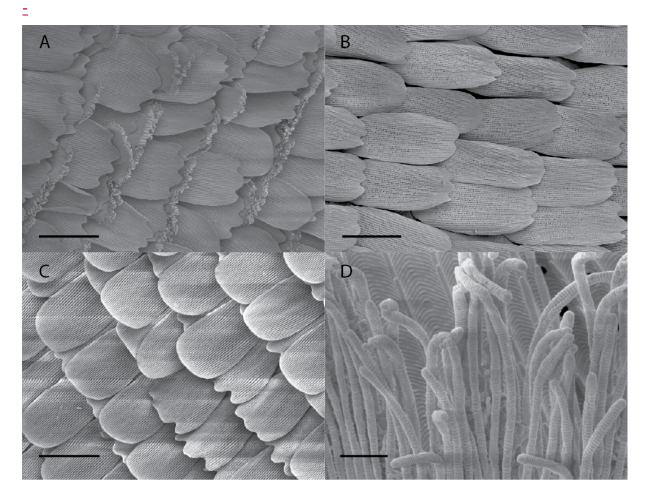
androconial region, with male (left) and female (right). C) Expanded view of the androconial

497 region with arrow highlighting the vein Sc+R1. The pale grey-brown region in the male wing

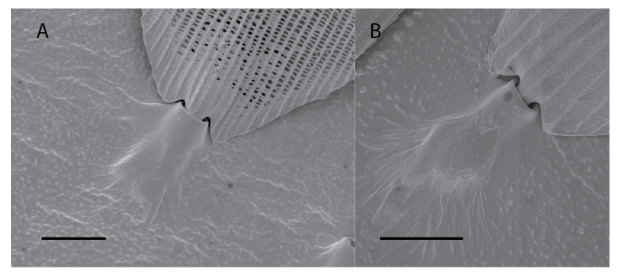
498 was dissected for chemical analysis.



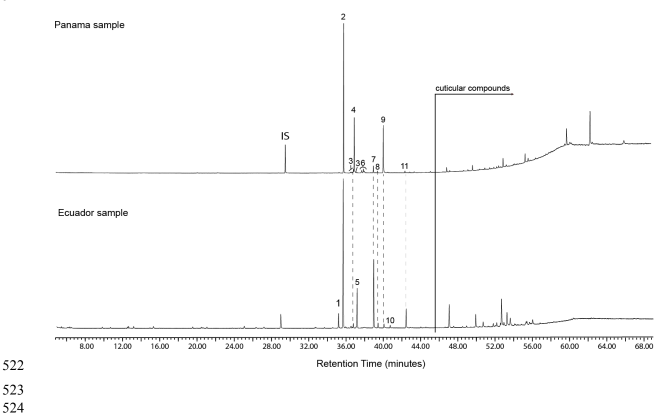
- 501 Figure 2: SEM image of putative androconial scales of *H. melpomene*. Scales in the region of
- 502 the hindwing vein Sc+R1 and forewing vein 1A are shown. (A) male hindwing (B) male
- 503 forewing and (C) female forewing at 500x magnification. (D) Magnified view of brush-like
- 504 structures of the special scales. Scale bars indicate 50 μm (A-C) and 2 μm (D).
- 505



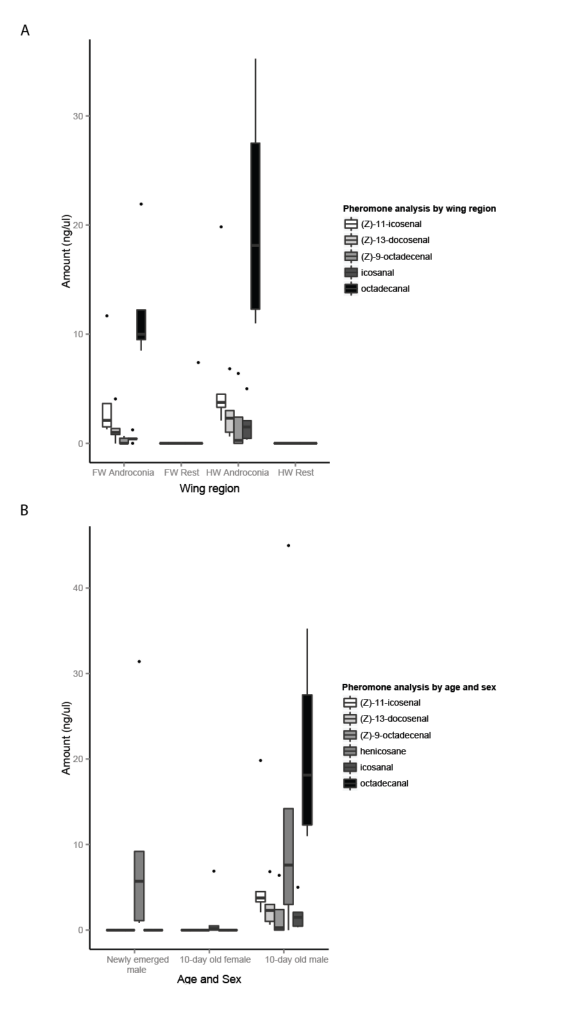
- 508 Figure 3: SEM image of the base of a scale of hindwing androconia of *H. melpomene* (A)
- scale from androconial region of male with brush-like structures. (B) scale from androconial
- 510 region of female. Scale bars indicate $10 \ \mu m$.

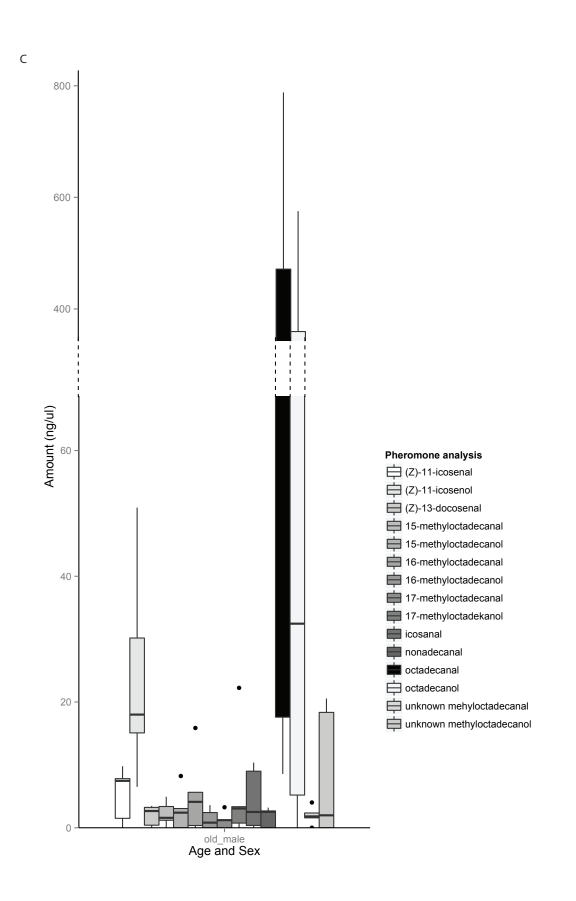


- Figure 4: Total ion chromatograms of extracts from the androconial region of an 513
- Ecuadorian *H. melpomene* and a Panamanian *H. melpomene* male hindwing. 1: (*Z*)-9-514
- octadecenal; 2: octadecanal; 3: 15-, 16-, 17-methyloctadecanals, and additional 515
- methyloctadecanal; 4: 1-octadecanol; 5: henicosane; 6: 15-, 16-, and 17-516
- methyloctadecan-1-ols, additional methyloctadecanol, and nonadecanal; 7: (Z)-11-517
- icosenal; 8: icosanal; 9: (Z)-11-icosenol; 10: tricosane; 11: (Z)-13-docosenal. All peaks 518
- eluting later than 44 min are cuticular compounds consisting of larger *n*-alkanes, 2,5-519
- dialkyltetrahydrofurans, cholesterol or are contaminations. 520
- 521



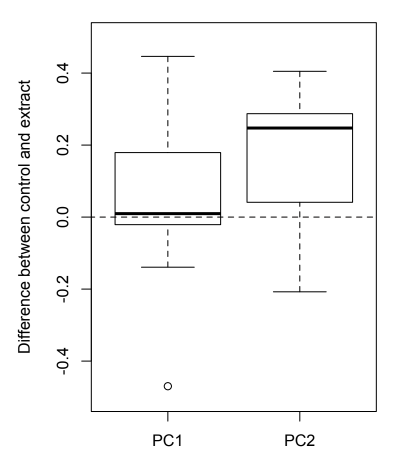
- 525 Figure 5: GC/MS analyses of *H. melpomene* wing extracts showing presence of putative male
- sex pheromone compounds (a) Presence of compounds in the hindwing androconial region of
- 527 five mature females, five young and five mature males, showing that compounds are largely
- restricted to mature males. (b) Presence of compounds in five mature females, five young and
- 529 five mature males, showing that compounds are largely restricted to mature males. (c)
- 530 Presence of compounds in hindwing androconial region of five wild males from Panama.







- 535 Figure 6: Differences in behavioural response for *H. melpomene* females presented with
- 536 models scented with male androconial extracts and unscented controls. Loading scores for
- 537 principal components are given in Table 1.



538 539

- 540 Table 1: Behavioural events scored for females presented with model butterflies and loading
- scores for *H. melpomene* female response to differential treatments. The mean proportion of
- 542 trials in which a given behavior was observed is shown for both Control and Extract models.
- 543 The full data set is given as a supplementary file.
- 544

Observed behaviour	PC1	PC2	Ctrl	Extract
Slow and moderate wing	0.15	-0.08	0.92	0.87
flapping				
Flying/Flying facing towards	-0.62	-0.12	0.45	0.41
model				
Slow rhythmic flight	-0.41	-0.26	0.23	0.31
Wing display with abdomen	0.25	-0.63	0.09	0.15
normal				
Quick and jerky wing flapping	0.60	0.11	0.60	0.53
Flying away from model	-0.05	0.11	0.98	0.96
Fast erratic flight	-0.09	0.70	0.35	0.19
Wings open with abdomen	-0.03	0.09	0.04	0.04
erect				