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1 Ancestral hymenopteran queen pheromones do not share the broad
2 phylogenetic repressive effects of honeybee queen mandibular
3 pheromone.

4

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24

25

26 Abstract

27 Queen pheromones effect the reproductive division of labour, a defining feature of
28 eusociality. Reproductive division of labour ensures that one, or a small number of, females
29 are responsible for the majority of reproduction within a colony. Much work on the
30 evolution and function of these pheromones has focussed on Queen Mandibular Pheromone
31 (QMP) which is produced by the Western or European honeybee (*Apis mellifera*). QMP has
32 phylogenetically broad effects, repressing reproduction in a variety of arthropods, including
33 those distantly related to the honeybee such as the fruit fly *Drosophila melanogaster*. QMP
34 is highly derived and has little chemical similarity to the majority of hymenopteran queen
35 pheromones which are derived from cuticular hydrocarbons. This raises the question of
36 whether the phylogenetically widespread repression of reproduction by QMP also occurs
37 with more basal saturated hydrocarbon-based queen-pheromones. Using *D. melanogaster* we
38 show that saturated hydrocarbons, are incapable of repressing reproduction, unlike QMP. We
39 also show no interaction between the four saturated hydrocarbons tested or between the
40 saturated hydrocarbons and QMP, implying that there is no conservation in the mechanism of
41 detection or action between these compounds. We propose that the phylogenetically broad
42 reproductive repression seen in response to QMP is not a feature of all queen pheromones,
43 but unique to QMP itself, which has implications for our understanding of how queen
44 pheromones act and evolve.

45 Introduction

46 Reproductive division of labour is a key feature of social insect societies, requiring a small
47 number of females, indeed often a single female, to be reproductively dominant, and her
48 subordinate workers to have their reproduction repressed (Oster and Wilson, 1978). To
49 achieve this in the Hymenoptera, a clade containing many eusocial species, a mixture of
50 behavioural aggression and chemical inhibition of reproduction is used (Le Conte and Hefetz,
51 2008; Padilla et al., 2016). Chemical inhibition occurs via queen pheromones (Matsuura et
52 al., 2010; Vargo and Laurel, 1994; Winston and Slessor, 1992). These queen pheromones are
53 produced by the reproductively dominant female and are thought to signal her fecundity to
54 subordinates (Keller and Nonacs, 1993). Queen pheromones have been thought to be
55 complex- both in function and composition (Brockmann et al., 1998; Slessor et al., 1988).
56 They are theorised to be the product of an evolutionary arms race between the dominant
57 female repressing reproduction, and the subordinate attempting to escape that repression (Le

58 Conte and Hefetz, 2008; Symonds and Elgar, 2008). This ‘escape’ could be achieved through
59 behavioural alterations or genetic changes that overcome reproductive repression, for
60 example through decreased sensitivity to or avoidance of the queen pheromone. It is
61 hypothesised that this may lead to the evolution of increasingly more elaborate pheromones
62 or mechanisms of repression, to accomplish reproductive dominance and eusociality (Katzav-
63 Gozansky, 2006).

64 Cuticular hydrocarbons (CHCs) are produced in the exocrine glands of insects, and secreted
65 into the cuticle (Howard and Blomquist, 2005). CHCs have various functions including
66 acting as contact pheromones to convey information both between and within species. CHCs
67 vary by species, sex, genotype, behavioural status, group within a society, reproductive state
68 and physiology in both solitary and social insects (Howard and Blomquist, 2005). Queen
69 signals within the social Hymenoptera are commonly CHCs, particularly long-chain linear
70 and methyl branched alkanes, although the nature of these compounds varies between species
71 (Van Oystaeyen et al., 2014). Van Oystaeyen *et al.* showed that saturated hydrocarbons
72 may act as sterility-inducing cues, or as indicators of fertility, in 57 out of 64 social
73 Hymenoptera, implying evolutionary conservation in the class of compound used to control
74 reproduction. This is remarkable as it indicates that CHCs have repeatedly become co-opted
75 into maintaining worker sterility in five independent origins of eusociality. The widespread
76 use of saturated hydrocarbons implies that these molecules may have functioned as fertility
77 cues in the ancestor of extant bees, ants and wasps (Van Oystaeyen et al., 2014). A subset of
78 these saturated hydrocarbons were the linear alkanes; pentacosane ($n\text{-C}_{25}$), heptacosane ($n\text{-}$
79 C_{27}), octacosane ($n\text{-C}_{28}$) and nonacosane ($n\text{-C}_{29}$) which were identified in a bumblebee, a
80 wasp and an ant species (*Bombus terrestris*, *Vespula vulgaris* and *Cataglyphis iberica*).
81 These compounds were functionally demonstrated to act as sterility – inducing queen
82 pheromones (Van Oystaeyen et al., 2014). It is important to note, however, that $n\text{-C}_{25}$ is found
83 on the cuticle, as well as in most exocrine glands of both reproductive and non-reproductive
84 bumble bees (Amsalem et al., 2014; Amsalem et al., 2009) which is not consistent with it
85 functioning as a queen pheromone. No single linear alkane molecule repressed reproduction
86 in all three species, however, all tested species had their reproductive capacity reduced by at
87 least one of these compounds. This is consistent with the idea that linear alkanes in particular
88 may have convergently evolved a role as queen pheromones in the eusocial Hymenoptera. It
89 is proposed these signalling molecules may have evolved from chemical cues of a solitary
90 ancestor (Van Oystaeyen et al., 2014) ~180 million years ago (Peters et al., 2017).

91 Despite the importance of saturated hydrocarbons as hymenopteran queen pheromones,
92 Queen Mandibular Pheromone (QMP), produced by queen honeybees (*Apis mellifera*), is still
93 the most studied social insect queen pheromone (Keeling et al., 2003; Pankiw et al., 1994;
94 Slessor et al., 1988). Although tergal gland secretions, which contain alkene hydrocarbons,
95 have been implicated in repressing worker reproduction (Wossler and Crewe, 1999) the effect
96 is generally much smaller than observed for QMP (Holman, 2018). QMP is, however,
97 highly derived and distinct from other social hymenopteran queen pheromones (Van
98 Oystaeyen et al., 2014). It is a complex pheromone with five major components - none of
99 which are saturated hydrocarbons. QMP has been shown to repress oogenesis in virgin *D.*
100 *melanogaster* females, producing ovaries with fewer mature oocytes when exposed
101 (Camiletti et al., 2013; Sannasi, 1969). This is surprising given the evolutionary distance
102 between honeybees and *Drosophila* is ~340 million years (Misof et al., 2014). QMP also
103 represses reproduction in a variety of other arthropods, including a species of ant (Carlisle
104 and Butler, 1956), termite (Hrdý et al., 1960), house fly (Nayar, 1963) and even a prawn
105 (Carlisle and Butler, 1956), spanning evolutionary distances of more than ~530 million years
106 (Misof et al., 2014).

107 We hypothesise that there are two possible scenarios for the broad phylogenetic range over
108 which QMP represses reproduction. The first is that QMP has evolved to target highly
109 conserved pathways to repress reproduction. This scenario is consistent with our current
110 molecular understanding of how QMP acts to control reproduction in the honeybee ovary
111 (Duncan et al., 2016; Ronai et al., 2016). These studies demonstrate that QMP modulates
112 highly conserved processes within the honeybee ovary (Duncan et al., 2016; Ronai et al.,
113 2016) and Notch signalling in particular is known to be environmentally responsive (Hsu and
114 Drummond-Barbosa, 2011). This may imply that QMP has evolved to target ancient
115 mechanisms for responding reproductively to environmental stimuli. QMP has evolved over
116 the last ~55 million years (Peters et al., 2017). However, despite evolving relatively recently
117 it is capable of repressing reproduction in species 530 million years diverged (Carlisle and
118 Butler, 1956; Misof et al., 2014). If QMP has evolved to modulate conserved pathways
119 controlling reproduction other, less derived, queen pheromones might not be expected to
120 share this effect. Therefore, the broad range of arthropods repressed by QMP would not be
121 similarly repressed by other social hymenopteran queen pheromones.

122 The second option is that all of the cuticular hydrocarbons, identified as putative basal queen
123 pheromones in the Hymenoptera, are capable of repressing reproduction in a phylogenetically

124 broad range of animals, similar to QMP. This would mean the repression of reproduction
125 observed by Van Oystaeyen *et al.* may be attributed to conserved mechanisms being targeted
126 by all hymenopteran queen pheromones, not only QMP. If *D. melanogaster* reproduction is
127 impaired by these ancestral-like queen pheromones then this would imply that this broader
128 class of compounds are also targeting conserved mechanisms of reproductive repression, and
129 that this property is not unique to QMP. If linear alkanes do not repress reproduction in
130 *D. melanogaster*, then this implies these queen pheromones are specialised to act in the insect
131 groups in which sociality evolved and as such have a narrower phylogenetic span than that of
132 QMP.

133 In this study, we test the ability of saturated hydrocarbons to repress reproduction in *D.*
134 *melanogaster*. We also test for synergistic interactions between the linear alkanes and also
135 with honeybee QMP. This has implications for the pathways through which they act. If they
136 disrupt or potentiate the action of each other, it would suggest the derived QMP shares
137 mechanisms of detection or action with the ancestral social hymenopteran pheromones,
138 further informing us about their evolutionary trajectories. If they were acting through the
139 same mechanisms to cause reproductive repression it would indicate that selective pressure or
140 drift has acted to change the inputs (pheromones) into this pathway. If they do not interact, it
141 would indicate that the mechanisms through which QMP is detected or acts are different from
142 those used by linear alkanes. We conclude that the wide phylogenetic span of arthropod
143 species on which the repressive function of QMP acts is a derived evolutionary novelty, not a
144 feature of the broader class of queen pheromones from which QMP evolved.

145 **Methods**

146 ***D. melanogaster* stocks and maintenance**

147 The Oregon-R modENCODE line (Stock #25211 from the Bloomington *Drosophila* stock
148 centre) was used for all *D. melanogaster* work in this study. Stocks were maintained at 25 °C
149 on a 12 h : 12 h light/dark cycle. Flies were raised on a sugar/yeast medium; of 3L dH₂O, 200
150 g organic cornmeal, 50 g brewer's yeast, 140 g sugar, 20 ml propionic acid and 15 ml 10%
151 methyl *p*-hydroxybenzoate in absolute ethanol.

152 **Virgin collection**

153 Only virgin female *D. melanogaster* were used for this study. These were anaesthetised with
154 CO₂, and observed under a GXM-XTL stereomicroscope (GT Vision, UK), with
155 phenotypically virgin females being isolated based upon the characteristics of enlarged

156 abdomens, pale colouration and presence of the meconium. Virgins were collected within one
157 hour of emergence, isolated with other virgin females, and stored at room temperature for 24
158 hours.

159 **Pheromone dilutions**

160 *QMP dilutions for concentration gradient*

161 Queen Mandibular Pheromone (QMP) from queen honeybees (*A. mellifera*) is quantified in
162 Queen equivalents (Qe). One Qe is the amount a mated queen will produce in a 24 hours
163 (Pankiw et al., 1996). QMP contains five major components (Slessor et al., 1988), that make
164 up 1 Qe for a European mated queen in the following amounts; 200µg 9-keto-(*E*)-2-
165 decenoic acid (ODA), 80µg 9-hydroxy-(*E*)-2-decenoic acid (9-HDA), 20µg methyl-*p*-
166 hydroxybenzoate (HOB), and 2 µg 4-hydroxy-3-methoxyphenylethanol (HVA) (Pankiw et
167 al., 1996). QMP (Intko Supply Ltd, Canada) was dissolved in absolute ethanol to the
168 concentrations of 3.25 Qe, 6.5 Qe, 13 Qe and 26 Qe, and stored at -20 °C until use.

169 *Linear alkane dilutions and mixtures*

170 The linear alkanes pentacosane (*n*-C₂₅), heptacosane (*n*-C₂₇), octacosane (*n*-C₂₈) and
171 nonacosane (*n*-C₂₉) were used based on experimentally determined doses identified in the
172 study by Van Oystaeyen et al., 2014. These compounds had been identified as components
173 queen signals, from the bumblebee, wasp and ant species *B. terrestris*, *V. vulgaris* and *C.*
174 *iberica*. The values previously calculated were used to determine 1 Qe in the Van Oystaeyen
175 study, this was based on the absolute amount present upon the queen's cuticle. 26 Qe was
176 used for the treatment in this study (based on high QMP doses in *D. melanogaster* exposure
177 studies (Camiletti et al., 2013). In order to maximise the chances of finding similar biological
178 effects of linear alkanes we treated *D. melanogaster* with levels of the linear alkanes 26 fold
179 higher than those found in relevant queens from Van Oystaeyen et al. (2014). One Qe of each
180 alkane for use in *D. melanogaster* was defined as the highest amount produced by one of the
181 three species in Van Oystaeyen et al. (2014) *B. terrestris*; 232.5 µg *n*-C₂₅, *V. vulgaris*; 118 µg
182 *n*- C₂₇, 6.1 µg *n*-C₂₈, 19 µg *n*-C₂₉. The alkane blend was a combination of all four linear
183 alkanes discussed each at 26 Qe. The linear alkanes were dissolved in HPLC grade pentane,
184 and stored at -20 °C. Pentane was used as the control treatment.

185 **Pheromone exposure in *D. melanogaster***

186 *QMP concentration gradient exposure in D. melanogaster*

187 Modified vials were made from 50 ml centrifuge tubes. Tubes were heated and the collection
188 end was removed. Two layers of Whatman number 1 filter paper shaped to fit the inside of

189 the lid and these were screwed into place. A cotton ball was used to plug to cut end of the
190 tube. Virgin *D. melanogaster* were aged for 24 hours, before being put in modified vials, and
191 500 µl of a liquid diet was added. This liquid diet was made fresh on the day of use in 5 ml
192 aliquots. The diet contains 4.75 ml dH₂O, 5% absolute ethanol, 0.15 g sugar and 0.1 g
193 brewer's yeast (Camiletti et al., 2013). On top of this diet, 20 µl of QMP solution was added.
194 The virgin *D. melanogaster* were anaesthetised with CO₂, and 10 were added to the vial lying
195 on its side, and allowed to recover before the vial was incubated upright at 25 °C for 48 hours.
196 Each treatment consisted of seven replicates and each replicate included 10 individuals (n =
197 70).

198 *Linear alkane exposure in D. melanogaster*

199 Diet and vial set up were as described for the concentration gradient above. On top of this
200 liquid diet, 100 µl of the linear alkane solutions were added. The virgin *D. melanogaster*
201 were anaesthetised with CO₂, and 10 were added to the vial lying on its side, and allowed to
202 recover before the vial was incubated upright at 25 °C for 48 hours. *D. melanogaster* were
203 exposed to each of the linear alkanes individually, as well as the mixture of all four. The
204 positive control for ovary repression was 26 Qe of *A. mellifera* QMP, dissolved in absolute
205 ethanol. Each treatment had five replicates of 10 individuals (n=50).

206 *Linear alkane and QMP combined exposure in D. melanogaster*

207 Virgin *D. melanogaster* were exposed to a combined exposure of linear alkanes and a low
208 dose of honeybee QMP. QMP was diluted to a dose of 3.25 Qe. This low dose was designed
209 to induce minor repression, allowing for further reduction in mature oocyte number should
210 the linear alkane mix interact synergistically interact with QMP. Ethanol was used as the
211 solvent control for QMP. There were three control combinations used to test the interaction
212 between the alkane mix and QMP; ethanol and pentane, linear alkane mix and ethanol, 3.25
213 Qe QMP and pentane. Exposure method and timing was carried out as described for single
214 linear alkanes. The only difference was the addition of two treatments to the top of the liquid
215 food, as opposed to the one addition described above.

216 ***D. melanogaster* ovary dissection and fixation**

217 Ovary dissections were performed using a GXM-XTL stereomicroscope (GT Vision, UK)
218 after *D. melanogaster* had been briefly anaesthetised with CO₂. Ovaries were dissected into a
219 petri dish containing ice-cold PBS. Any ovaries that were damaged or lost oocytes in the
220 dissection process were discarded. These were stored in 400 µl PBS on ice until all
221 dissections were complete (less than 30 min).

222 PBS was removed from the microcentrifuge tube containing dissected ovaries, down to 50 μ l.
223 To the tube added 900 μ l PBS and 4% formaldehyde. Ovaries were rocked at room
224 temperature for 10 minutes. Fixative was removed, and ovaries were washed four times with
225 PTx (PBS with 0.1% TritonX). Fixed ovaries were stored in 70 % ultrapure glycerol at 4 °C
226 in the dark until slide mounting. Ovaries were stored in glycerol for at least 24 hours before
227 being bridge-mounted for microscopy. The number of mature (vitellogenic) oocytes was
228 determined by manual counting under a GXM-XTL stereomicroscope (GT Vision, UK) and
229 was used as a measure of fecundity (King, 1970).

230 **Statistical analysis**

231 The number of mature oocytes per ovary in the *D. melanogaster* was analysed using R Studio
232 version 3. 5. 2. Assessment of whether the data fit a normal distribution was carried out using
233 a Shapiro-Wilk test, all data showed a non-normal distribution and so Generalised Linear
234 Mixed Models (GLMMs) with a Poisson error structure were used using lme4, in all cases
235 treatment was treated as a fixed effect and the slide number as a random factor. The maximal
236 model was simplified using Analysis of Deviance (AOD) to assess the effect of removing
237 terms. Where an effect of treatment was found, pairwise comparisons between treatments
238 were carried out using emmeans using a Tukey post-hoc test, to correct for multiple testing.
239 Effect sizes (Log odds) and 95% confidence intervals were calculated from the GLMMs
240 using R Studio version 3. 5. 2. (Supplementary Fig. 1).

241 **Results:**

242 ***D. melanogaster* are reproductively repressed by honeybee QMP in a dose-dependent**

243 **manner**

244 As previously reported (Camiletti et al., 2013) the number of mature oocytes in a *D.*
245 *melanogaster* ovary is decreased in a dose-dependent manner by exposure to QMP for 48 h
246 (Fig. 1) (AOD $\chi^2 = 56.142$, $df = 4$, $p = 1.87 \times 10^{-11}$). At the lowest dose of QMP tested (3.25
247 Qe) the number of mature oocytes was repressed by 36% (Ethanol mean = 17.56, 3.25 Qe
248 mean = 11.27 $p = 0.0140$). The highest exposure tested (26 Qe) reduced the number of
249 mature oocytes by 71% (Ethanol mean = 17.56, 26 Qe mean = 5.02 $p = < 0.001$) (Fig. 1).

250 ***D. melanogaster* are not reproductively repressed by putative basal hymenopteran**

251 **queen pheromones**

252 To determine whether *D. melanogaster* are reproductively repressed by the putative
253 conserved social insect queen signals, virgin females were exposed to the linear alkanes
254 pentacosane (n -C₂₅), heptacosane (n -C₂₇), octacosane (n -C₂₈) and nonacosane (n -C₂₉). The
255 26 Qe dose of QMP was included as a positive control for reproductive repression. QMP

256 induced the expected repression (AOD $\chi^2 = 52.597$, $df = 7$, $p = 4.26 \times 10^{-9}$) (Fig. 2), reducing
257 the number of mature oocytes by 59% (Ethanol mean = 8.14, 26 Qe mean = 3.36 $p = < 0.001$).
258 Note that this is a slightly lower magnitude of repression than observed in Fig. 1, likely due
259 to differences in protein sources used for diet preparation. Consistent with this the solvent
260 only controls in Fig. 1 have fewer mature oocytes (Fig. 1, Ethanol mean = 17.56) than the
261 solvent only controls in Fig. 2 (Fig. 2, Ethanol mean = 8.14).

262 The high dose QMP positive control (26 Qe) was the only significant reproductive repression
263 observed, and none of the single linear alkanes tested altered the number of mature oocytes
264 produced ($n\text{-C}_{25} p = 0.987$, $n\text{-C}_{27} p = 1.000$, $n\text{-C}_{28} p = 1.000$, $n\text{-C}_{29} p = 1.000$). To test whether
265 these compounds interact to repress reproduction we treated *D. melanogaster* with a blend of
266 all four linear alkanes. This blend also did not cause reproductive repression and did not alter
267 the number of mature oocytes produced ($p = 1.000$).

268 **Linear alkanes and honeybee QMP do not act synergistically**

269 To test whether there was any synergistic interaction between QMP and the linear alkanes,
270 the high dose alkane mix (26 Qe) was given as well as a low dose of QMP (3.25 Qe) (Fig. 3).
271 This dose of QMP was chosen to induce minor repression (Fig. 1), but not to the same extent
272 observed as a result of exposure to 26 Qe (Fig. 1). By inducing a small reduction in the
273 number of oocytes, we sought to observe synergistic or antagonistic effects on reproduction
274 between QMP and the linear alkanes.

275 There was no effect of any of these treatments on the number of mature oocytes in this
276 experiment (AOD $\chi^2 = 3.45$, $df = 3$, $p = 0.3273$) (Fig. 3). Consistent with Fig. 2 there was no
277 statistically significant repression induced by the high-dose alkane mix. The low-dose QMP
278 with pentane control acted as anticipated, where there was a small reduction in the number of
279 mature oocytes produced (decreasing the number of mature oocytes by ~23%, Pentane +
280 ethanol mean = 7.59, 3.25 Qe mean = 5.82), but this was not statistically significant.

281 Treatment with 3.25 Qe QMP and the blend of linear alkanes did not affect the number of
282 mature oocytes produced. This demonstrates that there is no synergistic interaction between
283 the linear alkanes and honeybee QMP and that QMP was not able to potentiate the effects of
284 these linear alkanes to cause reproductive repression.

285 **Discussion**

286 As in previous studies, we have shown that *D. melanogaster* are reproductively repressed by
287 honeybee QMP (Fig.1) (Camiletti et al., 2013). Initially, this appears an unusual phenomenon,

288 as *Drosophila* are not eusocial and are not closely related to honeybees (they are separated by
289 ~340 million years of evolution (Misof et al., 2014)). *Drosophila* would also be unlikely to
290 come into contact with QMP in their natural environment as these species occupy very
291 different habitats to the honeybee. As such, they presumably have not evolved to specifically
292 repress reproduction in response to QMP. Additionally, QMP mediated repression of
293 reproduction in non-target species is well established (Camiletti et al., 2013; Carlisle and
294 Butler, 1956; Hrdý et al., 1960; Nayar, 1963); this suggests QMP may have evolved to target
295 conserved pathways to repress reproduction. What remains unknown is the evolutionary
296 history of this response- namely, is this wide phylogenetic span of repression a feature that is
297 derived and novel to honeybee QMP? Or is it a feature of the ancestral queen pheromones
298 from which QMP has presumably evolved?

299 Linear alkanes have been identified to act as a conserved class of repressive cues in the
300 Hymenoptera by Van Oystayaen *et al.* In this study, we tested the ability of these queen
301 pheromones to repress reproduction in *D. melanogaster*. We show that the linear alkanes
302 pentacosane (*n*-C₂₅), heptacosane (*n*-C₂₇), octacosane (*n*-C₂₈) and nonacosane (*n*-C₂₉) do not
303 reduce the number of mature oocytes produced in the *D. melanogaster* ovary (Fig. 2). We
304 also tested the hypothesis that these compounds may interact additively or synergistically to
305 repress reproduction, but a blend of all four alkanes also showed no reduction in the number
306 of mature oocytes (Fig.2). Determining appropriate doses of individual queen-pheromones
307 to test (Holman et al., 2017), particularly in cross species comparisons as presented here, is
308 challenging. In this study we verified that maximal repression of ovary activity by QMP was
309 observed with 26 qe of QMP, similar to that previously reported (Camiletti et al., 2013).
310 Similarly, we chose to treat *Drosophila* with 26 qe of the individual linear alkanes,
311 calculating qe based on the highest levels of individual linear alkanes found in either *B.*
312 *terrestris*, *V. vulgaris* or *C. iberica* queens (Van Oystaeyen et al., 2014), rather than test
313 identical microgram quantities of each linear alkane. Our rationale was that these linear
314 alkanes are present in *B. terrestris*, *V. vulgaris* and *C. iberica* at different levels and that this
315 may reflect differences in biological activity of these compounds. To maximise the
316 likelihood of finding a physiological effect of these compounds, but remain within the realm
317 of physiologically relevant doses that workers of these species might be exposed to, we
318 treated *D. melanogaster* with doses of linear alkane 26 fold higher than produced by queens
319 of *B. terrestris*, *V. vulgaris* and *C. iberica*. It is also important to note that *D. melanogaster*
320 has a smaller biomass than either *B. terrestris* and *V. vulgaris* workers and is similar to *C.*

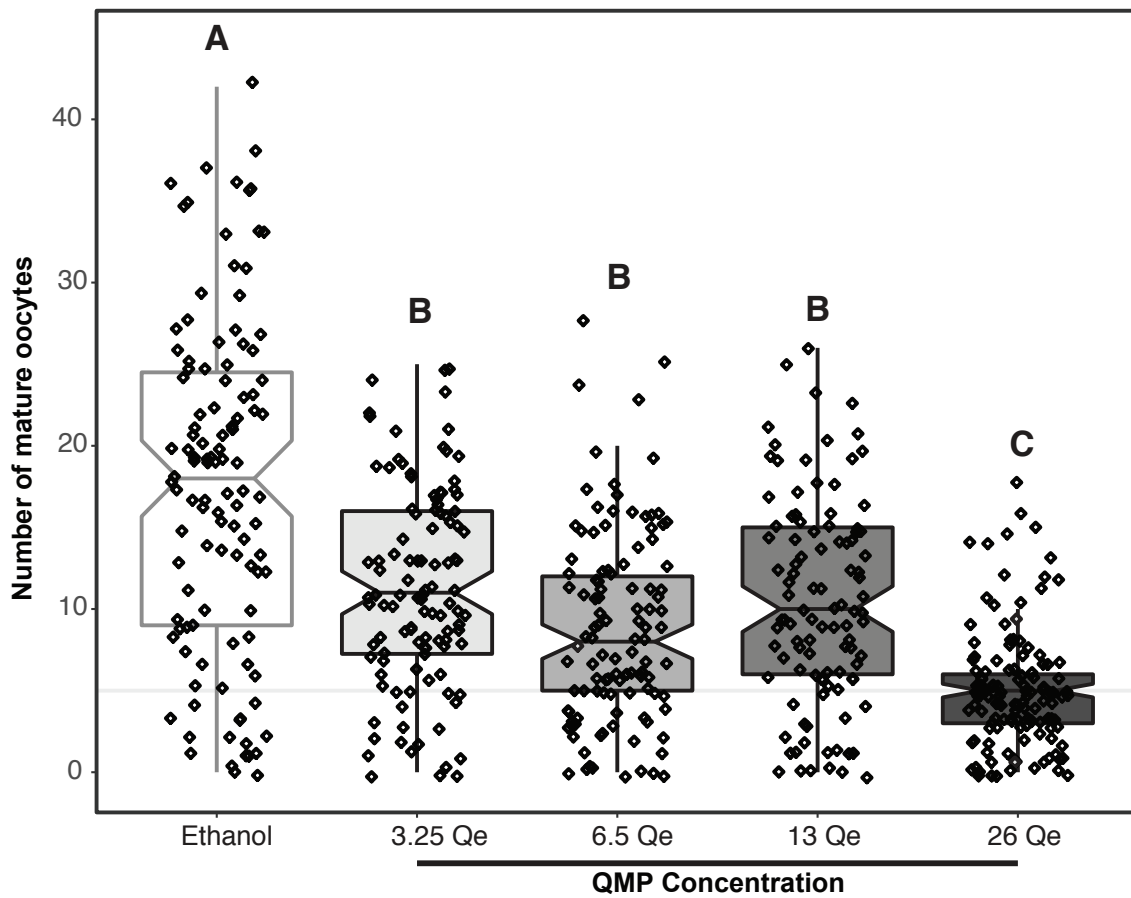
321 *iberica* so that the relative dose *D. melanogaster* were exposed to in this study is potentially
322 higher than 26 fold and higher than workers of these species would be exposed to. That *D.*
323 *melanogaster* don't respond to these relatively high doses of linear alkanes is consistent with
324 these compounds not having any biological activity in repressing ovary activity in *D.*
325 *melanogaster*.

326 That these basal Hymenopteran queen-pheromones don't affect reproduction in *D.*
327 *melanogaster* may be due to an inability of *D. melanogaster* to detect these compounds.
328 These linear alkanes tested in this study are derived from cuticular hydrocarbons (Van
329 Oystaeyen et al., 2014), which are known to vary in both quantity and identity between
330 species (Blomquist and Bagnères, 2010) so much so that even sister species can have distinct
331 cuticular hydrocarbon profiles (Morrison and Witte, 2011). It could be that these compounds
332 are unable to effect reproduction in *D. melanogaster* because they are not detected. However,
333 all four linear alkanes tested are components of the *D. melanogaster* CHC profile,
334 pentacosane (*n*-C₂₅) varies with geographical location in *D. melanogaster* populations
335 (Rajpurohit et al., 2017) and all four of the linear alkanes tested in this study vary in male *D.*
336 *melanogaster* with social group and genotype (Kent et al., 2008) suggesting that this species
337 can detect and respond these compounds. It is also possible that these compounds may be
338 affecting other aspects of *D. melanogaster* reproductive biology that were not examined in
339 this study, such as courtship or mating, as CHC profiles are known to be altered by mating
340 status (Everaerts et al., 2010). We hypothesise that rather than a lack of detection, these
341 compounds are targeting a pathway or process to repress reproduction that isn't conserved
342 between hymenoptera and *Drosophila*. This may be due to a loss of function of in the
343 lineage leading to *D. melanogaster* or a gain of function in the hymenopteran lineage. To test
344 this hypothesis we need a mechanistic understanding of how these linear alkanes are detected
345 and how this signal is translated into reproductive repression in social hymenoptera (Holman
346 et al., 2019).

347 We also tested whether there was a synergistic interaction between the linear alkanes and
348 QMP (Fig. 3). The linear alkane blend neither potentiated nor disrupted the minor repressive
349 effect of low dose QMP (Fig. 3). This implies that QMP is acting through a different
350 mechanism to the linear alkanes and as such can repress reproduction in *D. melanogaster*.
351 The response to QMP in *D. melanogaster*, and other non-target species, therefore is a derived
352 feature of QMP, not a reflection of a conserved class of 'insect pheromone'.

353 Surprisingly, QMP is capable of seemingly ubiquitously repressing highly diverged, non-
354 target species- whereas the other social hymenopteran queen pheromones cannot. QMP has
355 been evolving for ~ 55 million years (Peters et al., 2017), yet represses species that diverged
356 ~ 475 million years prior to the start of QMP evolving (Misof et al., 2014). This makes QMP
357 capable of targeting conserved pathways more ancient than QMP itself. The evolutionary
358 origins of QMP are unclear, but it has been hypothesised that an increase in social complexity
359 would be accompanied by an increase in the complexity of pheromones potentially as the
360 result of an arms race between queens and workers over worker reproduction where workers
361 evolve resistance to reproductive repression and queens evolve new pheromone components
362 that overcome that resistance (Bourke, 1988; Holman, 2018; Katzav-Gozansky, 2006; Kocher
363 and Grozinger, 2011). One consequence of such an arms race might be the evolution of
364 pheromones that target conserved pleiotropic pathways that are difficult to evolve resistance
365 to - as escape would come with high fitness costs. One such example is the previously
366 identified Notch cell signalling pathway (Duncan et al., 2016), which is key for QMP
367 mediated reproductive repression in honeybees but also has pleiotropic and conserved roles
368 in other fundamental processes including neurogenesis. These fundamental and pleiotropic
369 roles mean that there is selective pressure to retain a functioning Notch signalling pathway. If
370 such pathways are also evolutionary conserved, then targeting this system may cause
371 responses in a wide range of species- just as we see with QMP. Thus the broad effect of
372 QMP in arthropods is not a feature of ancestral queen pheromones, but instead may be a
373 derived feature unique to QMP itself. This may reflect that QMP has evolved to target an
374 evolutionarily conserved mechanism, possibly derived from an environmental signal linked
375 to temperature or nutrition, for repressing reproduction.

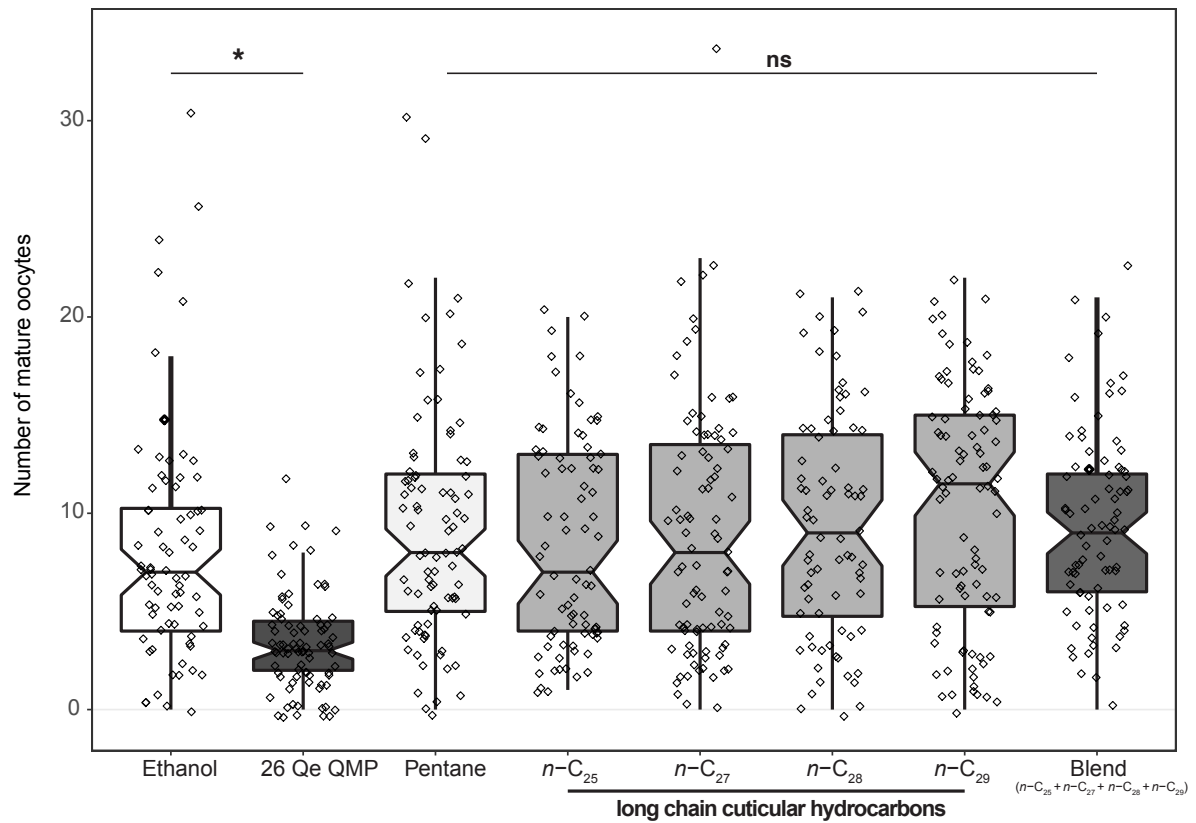
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378

379 **Figure 1.** A jittered box and whisker plot showing the number of mature oocytes from newly
 380 emerged virgin female *D. melanogaster* that were exposed to honeybee QMP in a
 381 concentration gradient from 3.25 – 26 Qe, with the ethanol solvent control. Exposure was for
 382 48 hours. Significant repression ($p < 0.05$) was induced at all concentrations tested relative to
 383 controls.

384



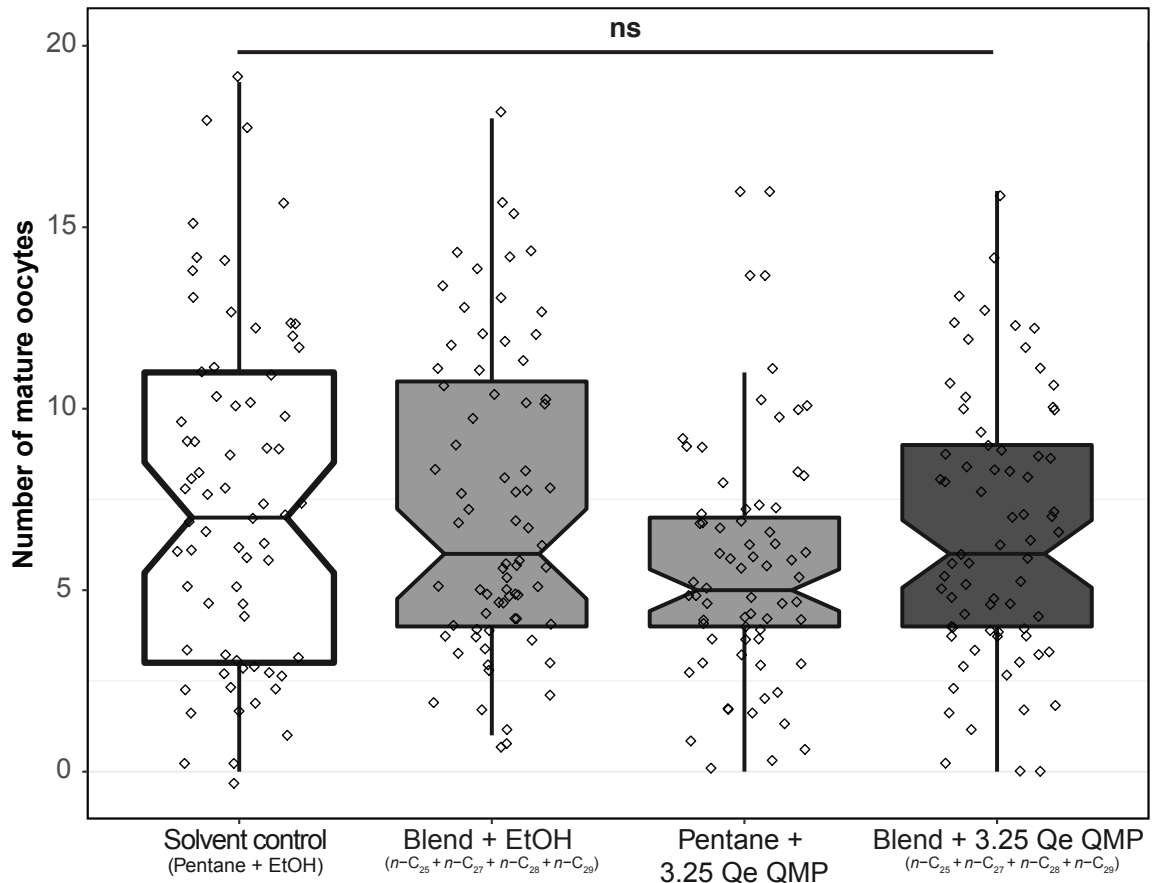
385

386 **Figure 2.** A jittered box and whisker plot showing the number of mature oocytes from newly
 387 emerged virgin female *D. melanogaster* that were exposed to the linear alkanes pentacosane
 388 ($n-C_{25}$), heptacosane ($n-C_{27}$), octacosane ($n-C_{28}$) and nonacosane ($n-C_{29}$) at dose of 26 Qe
 389 singularly, or as a blend of all four linear alkanes. Pentane was used as solvent control.
 390 Exposure was for 48 hours. 26 Qe of QMP from honeybees was used as a positive
 391 pheromone control, with the associated ethanol solvent control for QMP. The only
 392 statistically significant repression ($p < 0.05$) was induced by the high dose QMP.

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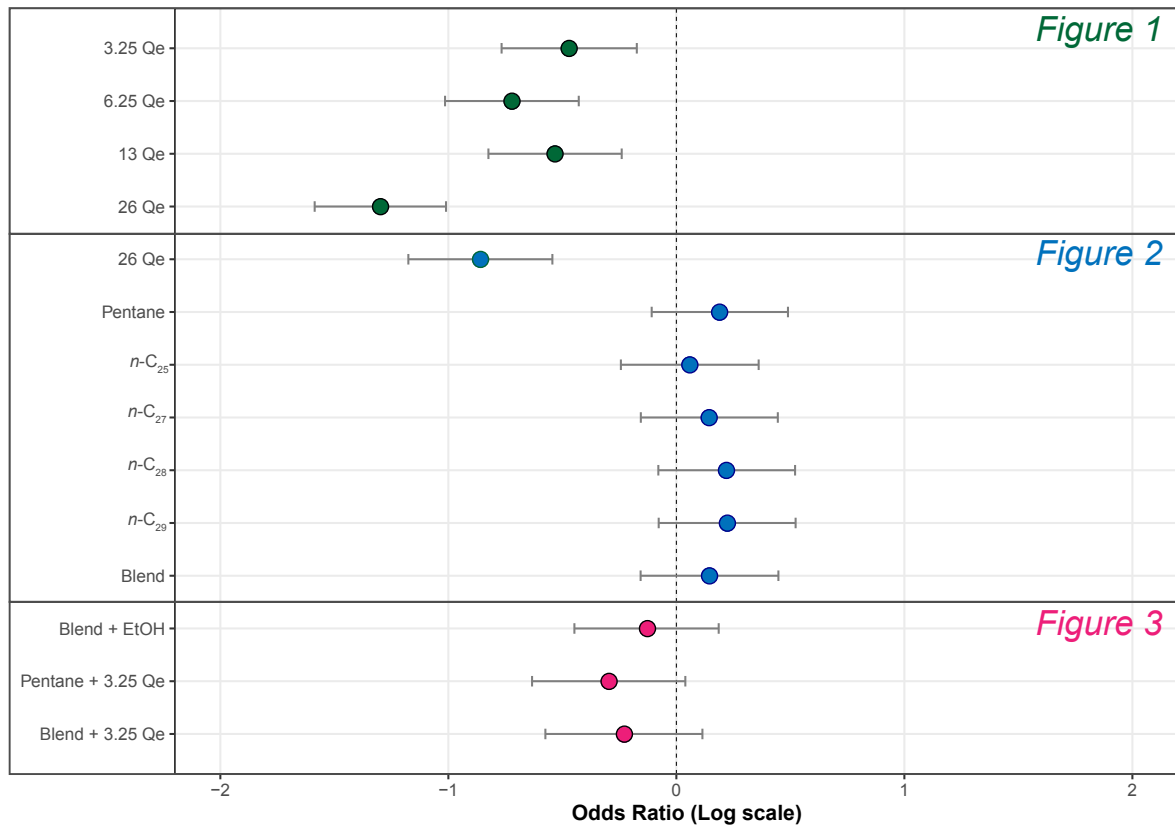
396

397 **Figure 3.**

398 A jittered box and whisker plot showing the number of mature oocytes from newly emerged
 399 virgin female *D. melanogaster* that were exposed to a blend of the four linear alkanes
 400 pentacosane ($n\text{-C}_{25}$), heptacosane ($n\text{-C}_{27}$), octacosane ($n\text{-C}_{28}$) and nonacosane ($n\text{-C}_{29}$) at dose
 401 of 26 Qe. Pentane was used as a solvent control for the alkanes. 3.25 Qe of QMP from
 402 honeybees was used to induce low levels of ovary repression, with the associated ethanol
 403 solvent control for QMP. Exposure was for 48 hours. There was no statistically significant
 404 repression induced by any of the treatments.

405

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408 **Supplementary Figure 1:** Effect sizes and 95% confidence intervals for data presented in
 409 Fig 1-3 of the main text. Effect sizes (Log odds) and 95% confidence intervals were
 410 calculated from the GLMMs using R Studio version 3. 5. 2. The only significant effects on *D.*
 411 *melanogaster* reproduction are for repression of reproduction by QMP (effect sizes do not
 412 overlap zero).

413

414

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422

423 **Author contributions**

424 MRL, PKD and EJD designed the study, MRL carried out the experimental work with
425 assistance from EJD. EJD and MRL performed the data analysis, prepared the figures and
426 drafted the manuscript with assistance from PKD. MRL, PKD, EJD edited and approved the
427 final manuscript.

428

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