Sunflower pollen reduces a gut pathogen in worker and queen but not male bumble bees

ALISON E. FOWLER, 1 © ELYSE C. STONE, 2 REBECCA E. IRWIN 3 © and LYNN S. ADLER 1 © 1 Department of Biology, University of Massachusetts Amherst, Amherst, Massachusetts, USA 2 Department of Biology, Smith College, Northampton, Massachusetts, USA and 3 Department of Applied Ecology, North Carolina State University, Raleigh, North Carolina, USA

Abstract. 1. Social insect castes and sexes differ in many ways, including morphology, behavior, and sometimes ploidy level. Recent studies have found that consuming sunflower pollen reduces the gut pathogen *Crithidia bombi* in workers of the common eastern bumble bee (*Bombus impatiens*). Here, this work is extended to the reproductive individuals that represent colony fitness – males and queens – to assess if the medicinal effects of sunflower pollen vary with bee caste and sex.

- 2. This study examined the effect of sunflower pollen compared to a diverse wildflower pollen mix on infection in worker, male, and daughter queen commercial *B. impatiens*. Bees were infected, fed either sunflower pollen or wildflower pollen for 7 days, and then infection levels were assessed.
- 3. Compared to wildflower pollen, sunflower pollen dramatically reduced *Crithidia* infection in workers and daughter queens, but not males. Infection levels were very low for both diets in males; this could be due to low pollen consumption or other mechanisms.
- 4. Reducing *Crithidia* infection in young queens before they undergo hibernation is important for population dynamics since infected queens are less likely to survive hibernation, and those that do are less likely to successfully establish a nest the following spring. Because sunflowers bloom in late summer when new queens are emerging, sunflowers could provide an important dietary component for queens during this critical life stage. Deepening our understanding of how diet impacts pathogens in reproductive bees, as well as workers, is crucial to maintain healthy pollinator populations.

Key words. *Bombus impatiens, Crithidia*, diet, pathogen resistance, pollinator, social caste.

Introduction

Individuals within a species may vary in pathogen loads due to factors such as sex, age, genotype, diet, social interactions, and experience. Females and males, specifically, may differ in traits related to pathogen defense due to differential exposure or impact of the pathogen on fitness (Rolff, 2002). In many invertebrates, including all members of the order Hymenoptera, differences between females and males are confounded by their different numbers of chromosomes; females are diploid, while males are typically haploid. Haploidy could negatively affect pathogen resistance by reducing the probability of inheriting resistance alleles (Mable & Otto, 1998; Otto &

Alison E. Fowler, 221 Morrill Science Center South, 611 North Pleasant St., Amherst, MA, 01003. Email: aefowler@umass.edu

Michalakis, 1998; O'Donnell & Beshers, 2004). However, evidence for this hypothesis has been equivocal, and differences between individuals might depend more on life history (Sheridan *et al.*, 2000). For example, one study compared male and female honey bees and paper wasps (Cappa *et al.*, 2015), which are both eusocial and haplodiploid, but males of the two species differ in how much they rely on protection from the colony. They found that paper wasp males, who spend less time in the colony, had higher immunocompetence than female workers, but honey bee males, who rely more heavily on colony protection, had lower immunocompetence than workers. This suggests that patterns of immunity are species-specific.

In social Hymenoptera, females develop into either reproductive queens or non-reproductive workers. Caste identity is determined by differential gene expression, often resulting in dramatic differences in morphology, behavior, longevity, and

physiology between queens and workers (Smith *et al.*, 2008). Given their longer lifespans, queens are expected to be equipped with higher immunocompetence than their non-reproductive sisters and daughters. This has been documented in honey bees (Chan *et al.*, 2006) and some ants (Gräff *et al.*, 2007; Koch *et al.*, 2013), but does not always result in lower infection in queens. For example, queen and worker honey bees had similar susceptibility to *Nosema ceranae* (Webster *et al.*, 2004), suggesting queens may evolve higher tolerance to certain pathogens rather than higher resistance. Understanding physiological and ecological differences between reproductive and non-reproductive females is important, especially since population dynamics in many species, such as bumble bees, are strongly influenced by queen survival and dispersal (Lepais *et al.*, 2010; Dreier *et al.*, 2014).

Like other social Hymenoptera, bumble bees (Bombus spp.) have queen, worker, and male individuals and are susceptible to a variety of pathogens, some of which have been implicated in their declines (e.g. Genersch et al., 2006; Cameron et al., 2011; Graystock et al., 2013a; Goulson et al., 2015). Worker bumble bees are the primary foragers and are recognised as important pollinators for both natural and agricultural ecosystems (e.g., Galen & Stanton, 1989; Elliott & Irwin, 2009; Garratt et al., 2014). Queen and male bumble bees are also important pollinators (Li et al., 2006; Ogilvie & Thomson, 2015) and represent colony fitness; yet pathogen dynamics in these individuals are poorly understood. Bumble bee queens had higher immune gene expression than males (Barribeau et al., 2015) and were less likely to acquire the gut pathogen, Crithidia sp., from their nest than workers (Ulrich et al., 2011). Males had lower infection levels than workers for several pathogens in the field (Shykoff & Schmid-Hempel, 1991) and had lower Crithidia infections than workers after 10 days in experimental trials (Ruiz-González & Brown, 2006), suggesting that males may have higher resistance than workers. Alternatively, adult male bumble bees do not spend as much time in the nest as workers and forage on different flowers (Roswell et al., 2019), which may reduce their chances of acquiring and transmitting pathogens (and thus may make them less favorable hosts). Improving our understanding of infection dynamics in queens and males is critical to conserve populations in the face of increasing pressure from pathogens (Meeus et al., 2011).

Differences in diet may also impact how individuals respond to pathogens. Host traits act within an environmental context, where factors such as food availability and intake can impact infection outcomes. Diet quantity and quality can impact host-pathogen interactions by either altering host condition or by directly feeding or inhibiting the pathogen. In the case of bumble bees infected with Crithidia, pollen starvation results in reduced immune gene expression (e.g., Brunner et al., 2014) but can also reduce infection levels (Logan et al., 2005; Conroy et al., 2016). Certain diets can also impact Crithidia infection. For example, infections decreased in bumble bees fed nectar with secondary compounds (e.g., Manson et al., 2010; Baracchi et al., 2015; Richardson et al., 2015), and infections were dramatically reduced in bees fed pollen from sunflowers (Helianthus annuus; Giacomini et al., 2018; LoCascio et al., 2019). These studies were all conducted on

female workers; how diet and pathogens interact in male and queen bumble bees is unknown. Males primarily forage for nectar, rather than pollen, resulting in different floral preferences and potentially lower pollen consumption compared to workers (Roswell *et al.*, 2019). As pollen starvation reduced *Crithidia* infection in bumble bee workers (Logan *et al.*, 2005; Conroy *et al.*, 2016), this might result in low infection levels in males if they eat little pollen. Queens may also respond to certain diets differently than workers given that they undergo different nutrient metabolism and storage (Colgan *et al.*, 2011).

The goals of this study were to examine the effect of pollen diet on pathogen infection in bumble bee queens, workers, and males. We experimentally infected commercial B. impatiens individuals with Crtihidia and asked how a diet of sunflower pollen affected infections compared to a wildflower pollen mix. We used male and worker bees in one experiment, and daughter queens in a separate experiment. This study asked two questions: (i) Do commercial B. impatiens males differ from workers in infection intensity when fed wildflower mix or sunflower pollen diet? and (ii) Does sunflower pollen reduce infection in commercial B. impatiens daughter queens? While we did not statistically compare queens to workers or males because they were studied in separate experiments, we discuss apparent patterns. Commercial bumble bees are used as pollinators for many outdoor and greenhouse crops and are susceptible to pathogens such as Crithidia (Graystock et al., 2013a). This work will improve our understanding of bee-diet-pathogen interactions, which may inform management strategies for bumble bee hives in agricultural settings, as well as conservation strategies for wild populations.

Materials and methods

Study system

Bombus impatiens (*Apidae*). The common eastern bumble bee is abundant in the eastern United States and currently not in decline (Cameron *et al.*, 2011). This species undergoes an annual life cycle; queens and males mate in late summer and then queens undergo solitary diapause during winter. Queens emerge in spring and initially lay worker eggs, progressing to males and daughter queens later in summer (Alford 1975). *Bombus impatiens* colonies are commercially available and are used commonly for crop pollination outdoors and in greenhouses.

Crithidia bombi (*Trypanosomatidae*). Crithidia bombi ("Crithidia" hereafter) is a gut pathogen of bumble bees transmitted horizontally through contact with infected feces on flowers or within the colony (Durrer & Schmid-Hempel, 1994; Otterstatter & Thomson, 2007). Crithidia directly reduces colony fitness; infected queens are 15% less likely to survive diapause (Fauser *et al.*, 2017), and infected queens that do survive are 40% less likely to successfully found a new colony (Brown *et al.*, 2003a). Infection is associated with decreased likelihood of reproduction in the wild (Goulson *et al.*, 2017), and infected workers have higher mortality when food-limited

(Brown et al., 2000). Previous studies have shown that Crithidia elicits an immune response in bumble bees (Brown et al., 2003b; Riddell et al., 2009), although this response is highly variable (Riddell et al., 2011; Brunner et al., 2013). It appears that the infection outcome is at least partly determined by the immune system given that increased immune gene expression of two antimicrobial peptides reduced Crithidia infection (Deshwal & Mallon, 2014). The gut microbiota, on the other hand, seems to play a major role in determining Crithidia infection; bees treated with antibiotics or reared in the absence of microbes suffered higher Crithidia loads than unmanipulated bees (Koch & Schmid-Hempel, 2011). Furthermore, infection was negatively correlated with the presence of certain bacterial taxa but positively correlated with bacterial diversity (Cariveau et al., 2014; Mockler et al., 2018; Näpflin & Schmid-Hempel, 2018). The success of infection is complex, and influenced by host genotype x pathogen genotype x microbiota genotype interactions (e.g. Barribeau & Schmid-Hempel, 2013; Näpflin & Schmid-Hempel, 2016).

Helianthus annuus (Asteraceae). Sunflower is a native US wildflower (Reagon & Snow, 2006) and major oilseed crop worldwide whose yield is improved by bee visitation (Nicolson & Human, 2013). In 2018, the United States planted 1.28 million acres of sunflowers (USDA Acreage Report, 2018). The medicinal effect of sunflower pollen in reducing Crithidia infection in B. impatiens was consistent across Crithidia strains (Giacomini et al., 2018) and sunflower cultivars (LoCascio et al., 2019). In addition, infection in wild-caught B. impatiens workers was negatively correlated with acreage of sunflowers (Giacomini et al., 2018). Sunflower pollen has relatively low protein content (Yang et al., 2013) and sometimes leads to poor performance in bees that feed on it (Tasei & Aupinel, 2008; McAulay & Forrest, 2019). The low protein content is not likely responsible for reducing Crithidia since another similarly low-protein pollen diet (buckwheat, Fagopyrum esculentum) resulted in comparatively high Crithidia infections (Giacomini et al., 2018). In addition, other chemical components of sunflower pollen, including several fatty acids and the secondary compounds rutin and triscoumaroyl spermidine derivatives, are not the mechanism by which sunflower pollen reduces Crithidia in the gut (Adler et al., 2020). Currently, the mechanism underlying this medicinal effect is unknown.

Experimental design

Two experiments were conducted, the first with workers and males and the second with commercial daughter queens. In both experiments, bees were inoculated with Crithidia, fed a pollen treatment for 7 days, and then dissected to assess pathogen cell counts. Half of each sex (worker/male) was randomly assigned to either wildflower pollen mix (Koppert Biological Systems, Howell, Michigan) or sunflower pollen (Helianthus annuus; Changge Hauding Wax Industry, China). The wildflower pollen mix (>10 plant species) represents a natural bumble bee diet and is also what commercial companies feed their hives and sell to customers and is therefore relevant

for both wild and commercial bees. The exact floral species composition is unknown; we purchased spring-collected pollen to avoid Asteraceae and checked the mix via microscopy for Asteraceae pollen (which is recognisable by its spiky exine) to ensure no sunflower or its relatives were in the mix. Goldenrod (Solidago spp.), another member of the Asteraceae family, was found to also have medicinal effects against Crithidia (LoCascio et al., 2019). The wildflower pollen mix is assumed to have a more diverse chemical and nutritional profile than sunflower pollen, and thus allows us to compare more typical dietary conditions of a generalist pollinator to a diet of only sunflower pollen. Because sunflower is relatively low in protein (Yang et al., 2013), we also suspect that the wildflower pollen mix is likely to have a higher protein concentration, although this was not tested. Both pollen types were collected by honey bees and were ground and mixed in a 7:1 ratio with 30% sucrose to make a paste that was frozen at -20 °C until use.

Upon entering the experiment, bees were inoculated with Crithidia originally from three wild B. impatiens workers from the Stone Soup Farm in 2014 (Hadley, Massachusetts: 42.363911 N, -72.567747 W) and then maintained in commercial colonies. Inoculum was prepared on each trial date with 150 µl of homogenised gut solution diluted with 1/4 strength Ringers Solution (Sigma Aldrich, St Louis, Missouri) to create a solution with 1200 cells/µl, which was then added to equal parts 50% sucrose solution for a final inoculum with 25% sucrose and 600 cells/µl. Bees were starved for 2 h, transferred to individual vials, presented with a 15-µl drop of inoculum (9000 pathogen cells, comparable to what bees would encounter in nature; Schmid-Hempel & Schmid-Hempel, 1993), and observed until the drop was consumed. Bees that did not consume the whole drop were excluded from the experiment.

After inoculation, bees were transferred into individual containers (Placon plastic deli cups with mesh bottoms and lids with holes; 11.4 cm top diameter, 8.9 cm bottom diameter, and 8.25 cm height) and administered their pollen treatment for 1 week. Bees were fed 10 ml of 30% sucrose along with 0.5 g of their pollen diet, replaced every other day. Bees were housed in an incubator in darkness at 27 °C and 55-60% humidity during the experiment. Crithidia infection reaches a representative population size by 7 days (Otterstatter & Thomson, 2006); thus, 7 days after inoculation, we dissected bees and assessed Crithidia cell counts. To dissect the gut, we removed the midgut and ileum and placed them in a 1.5-ml microcentrifuge tube with 300 µl of 1/4 strength Ringers Solution, homogenised, and left to settle for 4 h. We then placed a 10-µl aliquot of the supernatant on a hemocytometer (Hausser Scientific) and counted the number of moving Crithidia cells under a 400× compound light microscope to determine cells per 0.02 µl of gut solution. We measured marginal cell length of the right forewing of each bee to estimate bee size (Spaethe & Weidenmuller, 2002).

We did not measure pollen consumption in this experiment. Previous work has confirmed that workers and newly emerged queens consume pollen (Woodard et al., 2019) and that workers consume similar quantities of wildflower pollen mix and sunflower pollen (Adler et al., 2020). However, pollen consumption by male bumble bees has not been previously documented since male bees primarily forage for nectar during adulthood. To estimate the amount of pollen consumed by males, in a separate assay, we fed uninfected males and workers wildflower pollen for 3 days, dissected their guts following the protocol above, and counted pollen grains in 0.1- μ l of gut solution.

Bombus impatiens males and workers were sourced from three commercial colonies: two from BioBest LTD (Leamington, Ontario, Canada) and one from Koppert Biological Systems (Howell, Michigan). Oueens were sourced from six BioBest colonies; one colony provided individuals for both experiments. Each colony was checked for the absence of Crithidia upon arrival and bi-weekly thereafter by screening five workers. We used 79 males and 87 workers. Sixteen males and 14 workers died and were excluded from analyses, resulting in 63 males and 73 workers. We used 25 daughter queens; all survived their 7-day trial. All bees were removed from their natal colony at least 24h after emergence to allow inoculation of colony gut microbiota. The experiment was performed in spring and summer 2018 (File S1). For the pollen consumption trial, we used 24 males and 4 workers from one parent colony and 4 more workers from a separate parent colony. Data are available online in the Files S2-S4.

Statistical analyses

We used the open-source software R v3.3.3 (R Core Team, 2014) to analyse the number of Crithidia cells in a 0.02-µl gut extract as the response, with generalised linear mixed models (lme4 package, Bates et al., 2015). Cell counts were over-dispersed, and thus we used a negative binomial error distribution. We ran two analyses using the same general approach: one for workers/males and one for queens. For both models, the significance of terms was tested with likelihood ratio χ^2 tests (conducted with the ANOVA function in car package, Fox & Weisberg, 2019), which compares relative goodness of fit between models with and without each term. For all models, diet was included as a fixed effect. Initially, inoculation date and natal colony were included as random effects, but both were non-significant (P = 1 and 0.999, respectively, in worker/male model and P = 1 for both effects in queen model) and thus were excluded. We included wing marginal cell length as a covariate in each model because bee size can negatively correlate with Crithidia count (i.e., Richardson et al., 2015). For males and workers, bee sex (male vs. worker) was used as an additional fixed effect, along with bee sex by diet interaction. Although males were significantly larger than workers (F = 82.89, d.f. = 134, P < 0.0001), the Variance Inflation Factor indicated low multicollinearity in the model including both sex and wing marginal cell length (VIF < 2) and thus both were retained in the model. For the pollen consumption trial, bee sex (male vs. worker) was used as the fixed effect with pollen grain count as the response variable in a generalised linear model with a negative binomial error distribution. Bee wings were not collected for the pollen consumption trials, and thus, bee size was not accounted for in this analysis. Plots were made using emmeans (Lenth, 2020) and ggplot2 (Wickham, 2016). R code is available in the File S5.

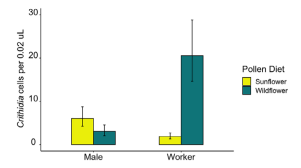


Figure 1. Effect of pollen diet on *Crithidia bombi* cell counts per 0.02 µl of gut solution in *Bombus impatiens* workers and males. Means estimated by a generalised linear model; error bars indicate standard error back-transformed by emmeans. [Colour figure can be viewed at wileyonlinelibrary.com].

Results

Crithidia cells replicated in hosts; when fed wildflower pollen, average raw counts in 0.02 µl of gut were between 2.74 and 22.08 cells, which is, on average, approximately 21 times more Crithidia cells than the initial inoculation. For males and workers, we found a significant interaction of diet and bee sex (male vs. worker) on *Crithidia* counts ($\chi^2 = 18.9831$, d.f. = 1, P < 0.0001; Fig. 1) and no significant relationship between wing cell length and infection ($\chi^2 = 0.6464$, d.f. = 1, P = 0.4214). After finding a significant interaction effect, we analysed each sex separately to determine responses to diet treatments. Workers fed wildflower pollen had over 10 times higher Crithidia counts than those fed sunflower ($\chi^2 = 23.2296$, d.f. = 1, P < 0.0001), but diet did not significantly affect Crithidia counts in males ($\chi^2 = 1.7909$, d.f. = 1, P = 0.1808; Fig. 1). Workers had over eight times higher Crithidia counts than males when fed wildflower pollen ($\chi^2 = 15.6267$, d.f. = 1, P < 0.0001), while males and workers did not differ when fed sunflower pollen ($\chi^2 = 2.2739$, d.f. = 1, P = 0.1316).

In daughter queens, sunflower pollen reduced *Crithidia* by 99.5% compared to wildflower pollen ($\chi^2 = 8.6996$, d.f. = 1, P = 0.0032; Fig. 2). Wing cell length was not significantly related to infection in queens ($\chi^2 = 0.4207$, d.f. = 1, P = 0.5166).

In the pollen consumption trial, pollen grain count in the gut was significantly lower in males than workers ($\chi^2 = 5.677$, d.f. = 1, P = 0.017). Male bumble bees had an average of 14 pollen grains per 0.1 µl of gut solution after 3 days, compared to an average of 122.75 pollen grains per 0.1 µl of gut solution in workers. However, one male had 313 pollen grains in 0.1 µl of gut solution; the remaining males had fewer than 10 grains. If we remove this outlier, males had an average of one pollen grain per gut sample, and the effect size of sex is more statistically significant ($\chi^2 = 75.364$, d.f. = 1, P < 0.0001).

Discussion

We found significant differences in how sunflower pollen affected pathogen counts in female queens and workers

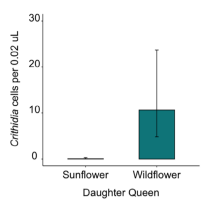


Figure 2. Effect of pollen diet on Crithidia bombi cell counts per 0.02 µl of gut solution in newly emerged Bombus impatiens daughter queens. Means estimated by a generalised linear model; error bars indicate standard error back-transformed by emmeans. [Colour figure can be viewed at wileyonlinelibrary.com].

compared to male bumble bees. Males responded differently to infection and diet than both castes of females. Although Crithidia grew in males relative to the initial inoculation dose, they exhibited remarkably low Crithidia counts compared to females regardless of diet, and there was no significant effect of diet on Crithidia counts. In workers and daughter queens, sunflower pollen dramatically reduced infection compared to wildflower pollen, consistent with previous findings (Giacomini et al., 2018; Adler et al., 2020).

Bumble bee queens are in a critical life stage when they first emerge. Before diapause, new queens have approximately 1 week in their natal nests before leaving to mate and overwinter (Goulson, 2010). It is during this critical window when queens sequester nutrients to prepare for diapause and when they are most likely to be exposed to pathogens. Prevalence of Crithidia increases over the season and is at its highest when daughter queens emerge (Popp et al., 2012). Crithidia persists from 1 year to the next by infecting queens before they overwinter, and therefore, there should be strong selection on the pathogen to infect queens. Since infected queens are 15% less likely to survive diapause (Fauser et al., 2017) and exhibit up to 40% lower colony-founding success than uninfected queens (Brown et al., 2003a), there should also be strong selection for queens to resist infection before entering into diapause. The effect of sunflower pollen on infection in daughter queens was dramatic; only 1 of the 13 queens fed sunflower pollen had any detectable Crithidia infection, and that was only one cell found in 0.02 µl of gut solution. Our study also corroborates findings that daughter queens were overall less likely to become infected with Crithidia than workers (Ulrich et al., 2011) since 7 of the 12 queens fed wildflower mix pollen had no detectable Crithidia infection.

As late-blooming flowers, sunflowers may be an important medicinal and nutritional food source for young queens in particular by increasing their chances of surviving diapause and founding a new colony in the spring. Recent research suggests that lipid and carbohydrate stores are essential for queen survival through diapause, whereas protein is not a critical macronutrient for storage (in contrast to its importance during larval development; Woodard et al., 2019). Sunflower pollen has relatively low protein and high lipid contents (Yang et al., 2013; Treanore et al., 2019). Therefore, sunflowers may provide new queens with important nutrients for surviving diapause, such as lipids, while also reducing Crithidia infections. Feeding on sunflower pollen during this life stage may therefore increase queen probability of surviving diapause and founding a colony in the spring (due to higher lipid stores and lower Crithidia infection). Queen survival through diapause constitutes significant population bottlenecks every year (Straub et al., 2015), and thus maximising macronutrient storage and minimising infection during the pre-diapause period is crucial for colony reproduction. Interestingly, Bombus impatiens workers are known to preferentially forage for pollen with high protein:lipid ratios to feed developing larvae (Vaudo et al., 2016); however, diet preferences in new adult queens before diapause may be different, or even reversed. Whether workers shift their foraging preferences when the colony starts to produce queens is an open area for future research.

Males had significantly lower baseline infection levels than workers (i.e. when fed the wildflower mix diet), and infection levels were not affected by diet. Previous work has found that male and worker bumble bees were similarly susceptible to Crithidia infection (Ruiz-González & Brown, 2006). This study also found that, for the first few days, males and workers had similar infection intensities, but over time, male infection intensities increased at a lower rate than workers, resulting in significantly lower cell counts in males by 10 days post-inoculation (Ruiz-González & Brown, 2006). These findings, along with our results, suggest that males do not suffer from haploid susceptibility and may, in fact, exhibit higher resistance to Crithidia than diploid workers. Alternatively, the low infections seen in males may be due to low pollen consumption since pollen starvation reduces Crithidia infection (Logan et al., 2005; Conroy et al., 2016). This hypothesis is supported by our pollen consumption trials, in which we found 10 times less pollen in the guts of male compared to worker bees. Nonetheless, studies comparing immune responses of males and workers would help clarify the mechanisms driving the observed patterns.

The bumble bee gut microbiota influences Crithidia infection (Koch & Schmid-Hempel, 2011; Koch & Schmid-Hempel, 2012, Cariveau et al., 2014; Mockler et al., 2018) and could be another mechanism driving differences between male and worker infection levels. Male and worker honey bees show differences in bacterial microbiota; males had lower bacterial diversity and higher prevalence of Lactobacillus "firm-5" in their guts than female workers (Kapheim et al., 2015). Lactobacillus "firm-5" abundance in the gut is negatively correlated with Crithidia infection in bumble bees, potentially by lowering the pH of the gut environment (Palmer-Young et al., 2018). If adult male bumble bees have higher prevalence of this bacteria than workers, that may be driving their low infection intensities.

The role of pollen nutritional content, including that of sunflower, in bee immune function is unclear. In bumble bees, immune gene expression is lower in bees starved of pollen, but immunocompetence was not affected by differences in dietary protein content in honey bees (Alaux et al., 2010) or bumble bees (Roger *et al.*, 2017). However, diet diversity may play an important role. Poly-floral diets increased immunocompetence in honey bees when compared to mono-floral diets (Alaux *et al.*, 2010). This suggests that sunflower pollen as a mono-floral diet (as used in our experiments) may reduce immune function in bees compared to a wildflower mixed diet. If sunflower pollen reduces immune function, then it likely inhibits *Crithidia* by another mechanism and may render bees susceptible to other infections that sunflower does not protect against. Future studies should focus on the specific roles that pollen diet, specifically sunflower pollen, plays in the bee immune system.

The mechanism underlying the medicinal effect of sunflower pollen is currently unknown. Chemical extracts from sunflower pollen reduced the growth of bacteria and fungi (Fatrcová-Šramková et al., 2016) but increased Crithidia growth in vitro (Palmer-Young & Thursfield, 2017) and did not reduce Crithidia to the level of pure sunflower pollen in bee assays (Adler et al., 2020). A compound in heather nectar (Calluna vulgaris) reduced Crithidia infection by removing the flagellum and preventing attachment to the gut wall (Koch et al., 2019); the spiky pollen coat of sunflower could mechanically inhibit attachment via similar mechanisms. Alternatively, sunflower pollen may influence Crithidia infection by inducing a community shift in the gut microbiome, an altered immune response, or changes in gut passage time; all of these mechanisms are currently being assessed in our research group. It is possible that these processes differ in male, queen, and worker bees. Adult males forage for nectar rather than pollen (Roswell et al., 2019), and new adult queens have different nutritional needs than developing larvae and adult workers (Vaudo et al., 2016; Woodard et al., 2019). These dietary differences may have large implications for the immune system and gut microbiota. Because the gut microbiota plays a role in resistance to Crithidia infection (e.g., Koch & Schmid-Hempel, 2011), caste and sex differences in the microbiota could affect how bees respond to both diet and pathogen infection.

Our results indicate that sunflower pollen effectively reduces a pathogen in commercial queens and workers and therefore could be used in commercial colony management. However, there are important distinctions between commercial and wild bumble bees to consider when interpreting these results in the context of wild populations. A previous study found that wild B. impatiens workers had higher susceptibility to Crithidia than commercially reared workers (Mockler et al., 2018), possibly due to the acquisition of non-core gut bacteria from the environment. In addition, commercial bees may be selected for resistance to pathogens that are often present in commercial rearing facilities (Graystock et al., 2013b). However, previous studies suggest that sunflower does reduce Crithidia in wild bumble bees (Giacomini et al., 2018) and that sunflower pollen is frequently collected and consumed by bumble bees and other pollinators (Westphal et al., 2003). These findings suggest that sunflower pollen, as part of a diverse diet, may be a promising natural remedy for bumble bee populations facing Crithidia infections. It is important to study both wild and commercial bees in order to best understand how they respond to diets and pathogens, especially since commercial and wild bees often share resources in agricultural settings (Graystock et al., 2016).

Queens and males are the reproductive individuals and so represent colony fitness, yet the vast majority of studies on bumble bees focus exclusively on workers. For example, worker bumble bees have been used to understand learning and behavior (Leadbeater & Chittka, 2007), the impacts of pesticides (Laycock et al., 2012), and immunity (Brown et al., 2003b). Meanwhile, males and queens are more difficult to acquire in large numbers, resulting in a knowledge gap about the ecology and physiology of individuals that will pass their genes to future generations. Pathogens are a strong selective force and an important factor contributing to wild and managed bee declines (Goulson et al., 2015); thus, it is crucial to understand how reproductive bees respond to pathogen infection to inform population conservation efforts. With a greater understanding of diet effects on bee pathogens, we could better manage pathogen spread by creating pollinator habitat tailored to provide bees with sufficient diet diversity and access to nutritional and medicinal food.

Acknowledgements

We thank N. Chevan, J. Cook, L. Coppinger, B. Joyce, A. McFarland, K. Michaud, and T. Shaya for research assistance; L. Katz for mentorship; Biobest and Koppert for donating bumble bee colonies; and members of the Adler lab and two anonymous reviewers for manuscript feedback. Research was funded by a Lotta Crabtree Fellowship from the University of Massachusetts College of Natural Sciences (AEF), National Science Foundation Graduate Research Fellowship (AEF), and USDA-NIFA-2016-07962 (LSA and REI). Any opinions, findings, and conclusions or recommendations expressed in this material are those of the authors and do not necessarily reflect the views of the funding agencies. The authors declare no conflict of interest.

AUTHOR CONTRIBUTION

LSA and REI designed the study, ECS and AEF collected the data, AEF analysed the data, and AEF wrote the paper with help from all authors.

Data availability statement

The data that supports the findings of this study are available in the Supporting Information.

Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

File S1 Table 1 with inoculation dates.

File S2 Daughter queens' data file.

File S3 Workers and males' data file.

File S4 Pollen grains' data file.

File S5 R script.

References

- AdlerLynn S., Fowler Alison E., Malfi Rosemary L., Anderson Patrick R., Coppinger Lily M., Deneen Pheobe M., Lopez Stephanie, Irwin Rebecca E., Farrell Iain W., Stevenson Philip C. (2020) Assessing Chemical Mechanisms Underlying the Effects of Sunflower Pollen on a Gut Pathogen in Bumble Bees. Journal of Chemical Ecology, http://dx.doi.org/10.1007/s10886-020-01168-4.
- Alaux, C., Ducloz, F., Crauser, D. & Le Conte, Y. (2010) Diet effects on honeybee immunocompetence. Biology Letters, 6, 562-565. https:// doi.org/10.1098/rsbl.2009.0986.
- Alford, D.V. (1975). Bumblebees. London: Davis-Poynter,
- Baracchi, D., Brown, M.J.F. & Chittka, L. (2015) Weak and contradictory effects of self-medication with nectar nicotine by parasitized bumblebees. F1000Research, 4, 1-15. https://doi.org/10 .12688/f1000research.6262.1.
- Barribeau, S.M., Sadd, B.M., du Plessis, L., Brown, M.J., Buechel, S.D., Cappelle, K. et al. (2015) A depauperate immune repertoire precedes evolution of sociality in bees. Genome Biology, 16, 83. https://doi.org/ 10.1186/s13059-015-0628-y.
- Barribeau, S.M. & Schmid-Hempel, P. (2013) Qualitatively different immune response of the bumblebee host, Bombus terrestris, to infection by different genotypes of the trypanosome gut parasite, Crithidia bombi. Infection, Genetics and Evolution, 20, 249-256. https://doi.org/10.1016/j.meegid.2013.09.014.
- Bates, D., Maechler, M., Bolker, B. & Walker, S. (2015) Fitting linear mixed-effects models using lme4. Journal of Statistical Software, 67, 1-48. https://doi.org/10.18637/iss.v067.i01.
- Brown, M.J.F., Moret, Y. & Schmid-Hempel, P. (2003a) Activation of host constitutive immune defence by an intestinal trypanosome parasite of bumble bees. Parasitology, 126, 253–260. https://doi.org/ 10.1017/S0031182002002755.
- Brown, M.J.F., Loosli, R. & Schmid-Hempel, P. (2000) Condition-dependant expression of virulence in trypanosome infecting bumblebees. Oikos, 91, 421-427.
- Brown, M.J.F., Schmid-Hempel, R. & Schmid-Hempel, P. (2003b) Strong context-dependent virulence in a host-parasite system: reconciling genetic evidence with theory. Journal of Animal Ecology, 72, 994 - 1002
- Brunner, F.S., Schmid-Hempel, P. & Barribeau, S.M. (2013) Immune gene expression in Bombus terrestris: signatures of infection despite strong variation among populations, colonies, and sister workers. PLoS One, 8, e68181. https://doi.org/10.1371/journal.pone.0068181.
- Brunner, F.S., Schmid-Hempel, P. & Barribeau, S.M. (2014) Protein-poor diet reduces host-specific immune gene expression in Bombus terrestris. Proceedings of the Royal Society B: Biological Sciences, 281, 20140128. https://doi.org/10.1098/rspb.2014.0128.
- Cameron, S.A., Lozier, J.D., Strange, J.P., Koch, J.B., Cordes, N., Solter, L.F. et al. (2011) Patterns of widespread decline in North American bumble bees. Proceedings of the National Academy of Sciences, 108, 662-667. https://doi.org/10.1073/pnas.1014743108.
- Cappa, F., Beani, L., Cervo, R., Grozinger, C. & Manfredini, F. (2015) Testing male immunocompetence in two hymenopterans with different levels of social organization: "live hard, die young?". Biological Journal of the Linnean Society, 114, 274-278. https://doi .org/10.1111/bij.12427.
- Cariveau, D.P., Powell, J.E., Koch, H., Winfree, R. & Moran, N.A. (2014) Variation in gut microbial communities and its association with pathogen infection in wild bumble bees (Bombus). The ISME Journal. 8, 2369-2379. https://doi.org/10.1038/ismej.2014.68.

- Chan, O.W.T., Howes, C.G. & Foster, L.J. (2006) Quantitative comparison of caste differences in honeybee Hemolymph. Molecular & Cellular Proteomics, 5, 2252–2262. https://doi.org/10.1074/mcp .M600197-MCP200.
- Colgan, T.J., Carolan, J.C., Bridgett, S.J., Sumner, S., Blaxter, M.L. & Brown, M.J.F. (2011) Polyphenism in social insects: insights from a transcriptome-wide analysis of gene expression in the life stages of the key pollinator, Bombus terrestris. BMC Genomics, 12, 1-21. https:// doi.org/10.1186/1471-2164-12-623.
- Conroy, T.J., Palmer-Young, E.C., Irwin, R.E. & Adler, L.S. (2016) Food limitation affects parasite load and survival of *Bombus impatiens* (Hymenoptera: Apidae) infected with Crithidia (Trypanosomatida: Trypanosomatidae). Environmental Entomology, 45, 1212-1219. https://doi.org/10.1093/ee/nvw099.
- Deshwal, S. & Mallon, E.B. (2014) Antimicrobial peptides play a functional role in bumblebee anti-trypanosome defense. Developmental and Comparative Immunology, 42, 240-243. https://doi.org/10.1016/ j.dci.2013.09.004.
- Dreier, S., Redhead, J.W., Warren, I.A., Bourke, A.F.G., Heard, M.S., Jordan, W.C. et al. (2014) Fine-scale spatial genetic structure of common and declining bumble bees across an agricultural landscape. Molecular Ecology, 23, 3384-3395. https://doi.org/10.1111/ mec.12823.
- Durrer, S. & Schmid-Hempel, P. (1994) Shared use of flowers leads to horizontal pathogen transmission. Proceedings of the Royal Society B: Biological Sciences, 258, 299-302. https://doi.org/10.1098/rspb .1994.0176.
- Elliott, S.E. & Irwin, R.E. (2009) Effects of flowering plant density on pollinator visitation, pollen receipt, and seed production in Delphinium barbeyi (Ranunculaceae). American Journal of Botany, 96, 912-919. https://doi.org/10.3732/ajb.0800260.
- Fatrcová-Šramková, K., Nôžková, J., Máriássyová, M. & Kačániová, M. (2016) Biologically active antimicrobial and antioxidant substances in the Helianthus annuus L. bee pollen. Journal of Environmental Science and Health Part B Pesticides, Food Contaminants, and Agricultural Wastes, 51, 176-181. https://doi.org/10.1080/03601234 .2015.1108811.
- Fauser, A., Sandrock, C., Neumann, P. & Sadd, B.M. (2017) Neonicotinoids override a parasite exposure impact on hibernation success of a key bumblebee pollinator. Ecological Entomology, 42, 306-314. https://doi.org/10.1111/een.12385.
- Fox, J., & Weisberg, S. (2019). An {R} Companion to Applied Regression, Third Edition. Thousand Oaks, CA: Sage Publications, URL https://socialsciences.mcmaster.ca/ifox/Books/Companion/
- Galen, C. & Stanton, M.L. (1989) Bumble bee pollination and floral morphology: factors influencing pollen dispersal in the alpine sky pilot, Polemonium viscosum (Polemoniaceae). American Journal of Botany, 76, 419-426. https://doi.org/10.1002/j.1537-2197.1989 .tb11330.x.
- Garratt, M.P.D., Coston, D.J., Truslove, C.L., Lappage, M.G., Polce, C., Dean, R. et al. (2014) The identity of crop pollinators helps target conservation for improved ecosystem services. Biological Conservation, 169, 128-135. https://doi.org/10.1016/j.biocon.2013
- Genersch, E., Yue, C., Fries, I. & De Miranda, J.R. (2006) Detection of deformed wing virus, a honey bee viral pathogen, in bumble bees (Bombus terrestris and Bombus pascuorum) with wing deformities. Journal of Invertebrate Pathology, 91, 61-63. https://doi.org/10 .1016/j.jip.2005.10.002.
- GiacominiJonathan J., Leslie Jessica, Tarpy David R., Palmer-Young Evan C., Irwin Rebecca E., Adler Lynn S. (2018) Medicinal value of sunflower pollen against bee pathogens. Scientific Reports, 8 (1), http://dx.doi.org/10.1038/s41598-018-32681-y.

- Goulson, D., Nicholls, E., Botias, C. & Rotheray, E.L. (2015) Bee declines driven by combined stress from parasites, pesticides, and lack of flowers. Science, 347, 1255957-1255957. https://doi.org/10.1126/ science.1255957.
- Goulson, D. (2010) Bumblebees: Behaviour, Ecology, and Conservation. 2, Oxford: Oxford University Press.
- Goulson, D., O'Connor, S. & Park, K.J. (2017) The impacts of predators and parasites on wild bumblebee colonies. Ecological Entomology., 43, 168-181. https://doi.org/10.1111/een.12482.
- Gräff, J., Jemielity, S., Parker, J.D., Parker, K.M. & Keller, L. (2007) Differential gene expression between adult queens and workers in the ant Lasius Niger. Molecular Ecology, 16, 675-683. https://doi.org/ 10.1111/j.1365-294X.2007.03162.x.
- Graystock, P., Blane, E.J., McFrederick, Q.S., Goulson, D. & Hughes, W.O.H. (2016) Do managed bees drive parasite spread and emergence in wild bees? International Journal for Parasitology: Parasites and Wildlife, 5, 64-75. https://doi.org/10.1016/j.ijppaw.2015.10.001.
- Graystock, P., Yates, K., Darvill, B., Goulson, D. & Hughes, W.O.H. (2013a) Emerging dangers: deadly effects of an emergent parasite in a new pollinator host. Journal of Invertebrate Pathology, 114, 114-119. https://doi.org/10.1016/j.jip.2013.06.005.
- Graystock, P., Yates, K., Evison, S.E.F., Darvill, B., Goulson, D. & Hughes, W.O.H. (2013b) The Trojan hives: pollinator pathogens, imported and distributed in bumblebee colonies. Journal of Applied Ecology, 50, 1207-1215. https://doi.org/10.1111/1365-2664.12134.
- Kapheim, K.M., Rao, V.D., Yeoman, C.J., Wilson, B.A., White, B.A., Goldenfeld, N. et al. (2015) Caste-specific differences in hindgut microbial communities of honey bees (Apis mellifera). PLoS One, 10, 1-14. https://doi.org/10.1371/journal.pone.0123911.
- Koch, H. & Schmid-Hempel, P. (2011) Socially transmitted gut microbiota protect bumble bees against an intestinal parasite. Proceedings of the National Academy of Sciences of the United States of America, 108, 19288-19292. https://doi.org/10.1073/pnas.1110474108.
- Koch, H., Woodward, J., Langat, M.K., Brown, M.J.F. & Stevenson, P.C. (2019) Flagellum removal by a nectar metabolite inhibits infectivity of a bumblebee parasite. Current Biology, 29, 3494-3500.e5. https:// doi.org/10.1016/j.cub.2019.08.037.
- KochHauke, Schmid-Hempel Paul (2012) Gut microbiota instead of host genotype drive the specificity in the interaction of a natural host-parasite system. Ecology Letters, 15, 1095-1103. http://dx.doi .org/10.1111/j.1461-0248.2012.01831.x.
- Koch, S.I., Groh, K., Vogel, H., Hannson, B.S., Kleineidam, C.J. & Grosse-Wilde, E. (2013) Caste-specific expression patterns of immune response and chemosensory related genes in the leaf-cutting ant, Atta vollenweideri. PLoS One, 8, 1-17. https://doi.org/10.1371/ journal.pone.0081518.
- Laycock, I., Lenthall, K.M., Barratt, A.T. & Cresswell, J.E. (2012) Effects of imidacloprid, a neonicotinoid pesticide, on reproduction in worker bumble bees (Bombus terrestris). Ecotoxicology, 21, 1937-1945. https://doi.org/10.1007/s10646-012-0927-y.
- Leadbeater, E. & Chittka, L. (2007) The dynamics of social learning in an insect model, the bumblebee (Bombus terrestris). Behavioral Ecology and Sociobiology, 61, 1789-1796. https://doi.org/10.1007/ s00265-007-0412-4.
- Lenth, R. (2020). emmeans: Estimated Marginal Means, aka Least-Squares Means. URL https://CRAN.R-project.org/ package=emmeans
- Lepais, O., Darvill, B., O'Connor, S., Osborne, J.L., Sanderson, R.A., Cussans, J. et al. (2010) Estimation of bumblebee queen dispersal distances using sibship reconstruction method. Molecular Ecology, 19, 819-831. https://doi.org/10.1111/j.1365-294X.2009.04500.x.
- Li, P., Luo, Y.B., Bernhardt, P., Yang, X.Q. & Kou, Y. (2006) Deceptive pollination of the Lady's Slipper Cypripedium tibeticum

- (Orchidaceae). Plant Systematics and Evolution, 262, 53-63. https:// doi.org/10.1007/s00606-006-0456-3.
- LoCascio, G.M., Aguirre, L., Irwin, R.E. & Adler, L.S. (2019) Pollen from multiple sunflower cultivars and species reduces a common bumblebee gut pathogen. Royal Society Open Science, 6, 190279. https://doi.org/10.1098/rsos.190279.
- Logan, A., Ruiz-González, M.X. & Brown, M.J.F. (2005) The impact of host starvation on parasite development and population dynamics in an intestinal trypanosome parasite of bumble bees. Parasitology, 130, 637-642. https://doi.org/10.1017/S0031182005007304.
- Mable, B.K. & Otto, S.P. (1998) The evolution of life cycles with haploid and diploid phases. BioEssays, 20, 453-462. https://doi.org/10.1002/ (SICI)1521-1878(199806)20:6<453::AID-BIES3>3.0.CO;2-N.
- Manson, J.S., Otterstatter, M.C. & Thomson, J.D. (2010) Consumption of a nectar alkaloid reduces pathogen load in bumble bees. Oecologia, 162, 81-89. https://doi.org/10.1007/s00442-009-1431-9.
- McAulay, M.K. & Forrest, J.R.K. (2019) How do sunflower pollen mixtures affect survival of queenless microcolonies of bumblebees (Bombus impatiens)? Arthropod-Plant Interactions, 13, 517-529. https://doi.org/10.1007/s11829-018-9664-3.
- Meeus, I., Brown, M.J.F., De Graaf, D.C. & Smagghe, G. (2011) Effects of invasive parasites on bumble bee declines. Conservation Biology, 25, 662-671. https://doi.org/10.1111/j.1523-1739.2011.01707.x.
- Mockler, B.K., Kwong, W.K., Moran, N.A., Koch, H. & Drake, H.L. (2018) Microbiome structure influences infection by the parasite Crithidia bombi in bumble bees. Applied and Environmental Microbiology, 84, 2335-2352. https://doi.org/10.1128/AEM.
- Näpflin, K. & Schmid-Hempel, P. (2016) Immune response and gut microbial community structure in bumblebees after microbiota transplants. Proceedings of the Royal Society B: Biological Sciences, 283, 20160312. https://doi.org/10.1098/rspb.2016.0312.
- Näpflin, K. & Schmid-Hempel, P. (2018) High gut microbiota diversity provides lower resistance against infection by an intestinal parasite in bumblebees. American Naturalist, 192, 131-141. https://doi.org/10 .1086/698013.
- Nicolson, S.W. & Human, H. (2013) Chemical composition of the "low quality" pollen of sunflower (Helianthus annuus, Asteraceae). Apidologie, 44, 144-152. https://doi.org/10.1007/s13592-012-0166-
- O'Donnell, S. & Beshers, S.N. (2004) The role of male disease susceptibility in the evolution of haplodiploid insect societies. Proceedings of the Royal Society B: Biological Sciences, 271, 979-983. https://doi .org/10.1098/rspb.2004.2685.
- Ogilvie, J.E. & Thomson, J.D. (2015) Male bumble bees are important pollinators of a late-blooming plant. Arthropod-Plant Interactions, 9, 205-213. https://doi.org/10.1007/s11829-015-9368-x.
- Otterstatter, M.C. & Thomson, J.D. (2006) Within-host dynamics of an intestinal pathogen of bumble bees. *Parasitology*, **133**, 749–761. https://doi.org/10.1017/S003118200600120X.
- Otterstatter, M.C. & Thomson, J.D. (2007) Contact networks and transmission of an intestinal pathogen in bumble bee (Bombus impatiens) colonies. Oecologia, 154, 411-421. https://doi.org/10 .1007/s00442-007-0834-8.
- Otto, S.P. & Michalakis, Y. (1998) The evolution of recombination in changing environments. Trends in Ecology and Evolution, 13, 145-151. https://doi.org/10.1016/S0169-5347(97)01260-3.
- Palmer-Young, E.C., Raffel, T.R. & McFrederick, Q.S. (2018) pH-mediated inhibition of a bumblebee parasite by an intestinal symbiont. Proceedings of the Royal Society B: Biological Sciences, 285, 1-9. https://doi.org/10.1098/rspb.2018.2041.
- Palmer-Young, E.C. & Thursfield, L. (2017) Pollen extracts and constituent sugars increase growth of a trypanosomatid parasite of bumble bees. PeerJ, 5, e3297. https://doi.org/10.7717/peerj.3297.

- Popp, M., Erler, S. & Lattorff, H.M.G. (2012) Seasonal variability of prevalence and occurrence of multiple infections shape the population structure of Crithidia bombi, an intestinal parasite of bumblebees (Bombus spp.). MicrobiologyOpen, 1, 362-372. https://doi.org/10 .1002/mbo3.35.
- Reagon, M. & Snow, A.A. (2006) Cultivated Helianthus annuus (Asteraceae) volunteers as a genetic "bridge" to weedy sunflower populations in North America. American Journal of Botany, 93, 127-133. https://doi.org/10.3732/ajb.93.1.127.
- Richardson, L.L., Adler, L.S., Leonard, A.S., Henry, K., Anthony, W.E., Manson, J.S. et al. (2015) Secondary metabolites in floral nectar reduce parasite infections in bumble bees. Proceedings of the Royal Society B: Biological Sciences, 282, 20142471. https://doi.org/10 .1098/rspb.2014.2471.
- Riddell, C., Adams, S., Schmid-Hempel, P. & Mallon, E.B. (2009) Differential expression of immune defences is associated with specific host-parasite interactions in insects. PLoS One, 4, 2-5. https://doi .org/10.1371/journal.pone.0007621.
- Riddell, C.E., Sumner, S., Adams, S. & Mallon, E.B. (2011) Pathways to immunit: temporal dynamics of the bumblebee (Bombus terrestris) immune response against a trypanosomal gut parasite. Insect Molecular Biology, 20, 529-540. https://doi.org/10.1111/j.1365-2583.2011 .01084.x.
- Roger, N., Michez, D., Wattiez, R., Sheridan, C. & Vanderplanck, M. (2017) Diet effects on bumblebee health. Journal of Insect Physiology, 96, 128-133. https://doi.org/10.1016/j.jinsphys.2016.11 .002.
- Rolff, J. (2002) Bateman's principle and immunity. Proceedings of the Royal Society B: Biological Sciences, 269, 867-872. https://doi.org/ 10.1098/rspb.2002.1959.
- RoswellMichael, Dushoff Jonathan, Winfree Rachael (2019) Male and female bees show large differences in floral preference. PLOS ONE, 14 (4), e0214909http://dx.doi.org/10.1371/journal.pone.0214909.
- Ruiz-González, M.X. & Brown, M.J.F. (2006) Males vs workers: testing the assumptions of the haploid susceptibility hypothesis in bumblebees. Behavioral Ecology and Sociobiology, 60, 501-509. https://doi.org/10.1007/s00265-006-0192-2.
- Schmid-Hempel, P. & Schmid-Hempel, R. (1993) Transmission of a pathogen in Bombus terrestris, with a note on division of labour in social insects. Behavioral Ecology and Sociobiology, 33, 319 - 327.
- Sheridan, L.A.D., Poulin, R., Ward, D.F. & Zuk, M. (2000) Sex differences in parasitic infections among arthropod hosts: is there a male bias? Oikos, 88, 327-334.
- Shykoff, J.A. & Schmid-Hempel, P. (1991) Incidence and effects of four parasites in natural populations of bumble bees in Switzerland. Apidologie, 22, 117-125. https://doi.org/10.1051/apido:19910204.
- Smith, C.R., Toth, A.L., Suarez, A.V. & Robinson, G.E. (2008) Genetic and genomic analyses of the division of labour in insect

- societies. Nature Reviews Genetics, 9, 735-748. https://doi.org/10 .1038/nrg2429.
- Spaethe, J. & Weidenmuller, A. (2002) Size variation and foraging rate in bumblebees (Bombus terrestris). Insectes Sociaux, 49, 142-146. https://doi.org/10.1007/s00040-002-8293-z.
- Straub, L., Williams, G.R., Pettis, J., Fries, I. & Neumann, P. (2015, December 1) Superorganism resilience: eusociality and susceptibility of ecosystem service providing insects to stressors. Current Opinion in Insect Science, 12, 109-112. https://doi.org/10.1016/j.cois.2015 .10.010.
- Tasei, J.-N. & Aupinel, P. (2008) Nutritive value of 15 single pollens and pollen mixes tested on larvae produced by bumblebee workers (Bombus terrestris, hymenoptera: Apidae). Apidologie, 39, 397–409. https://doi.org/10.1051/apido.
- Treanore, E.D., Vaudo, A.D., Grozinger, C.M. & Fleischer, S.J. (2019) Examining the nutritional value and effects of different floral resources in pumpkin agroecosystems on Bombus impatiens worker physiology. Apidologie, 50, 542-552. https://doi.org/10 .1007/s13592-019-00668-x.
- Ulrich, Y., Sadd, B.M. & Schmid-Hempel, P. (2011) Strain filtering and transmission of a mixed infection in a social insect. Journal of Evolutionary Biology, 24, 354–362. https://doi.org/10.1111/j.1420-9101.2010.02172.x.
- VaudoAnthony D., Patch Harland M., Mortensen David A., Tooker John F., Grozinger Christina M. (2016) Macronutrient ratios in pollen shape bumble bee (Bombus impatiens) foraging strategies and floral preferences. Proceedings of the National Academy of Sciences, 113 (28), E4035-E4042. http://dx.doi.org/10.1073/pnas.1606101113.
- Webster, T.C., Pomper, K.W., Hunt, G., Thacker, E.M. & Jones, S.C. (2004) Nosema apis infection in worker and queen Apis mellifera. Apidologie, 35, 49-54.
- Westphal, C., Steffan-Dewenter, I. & Tscharntke, T. (2003) Mass flowering crops enhance pollinator densities at a landscape scale. Ecology Letters, 6, 961-965. https://doi.org/10.1046/j.1461-0248 .2003.00523.x.
- Wickham, H. (2016). ggplot: Elegant Graphics for Data Analysis. New York: Springer-Verlag, URL https://ggplot2.tidyverse.org
- WoodardS Hollis, Duennes Michelle A, Watrous Kristal M, Jha Shalene (2019) Diet and nutritional status during early adult life have immediate and persistent effects on queen bumble bees. Conservation Physiology, 7, http://dx.doi.org/10.1093/conphys/coz048.
- Yang, K., Dan, W., Xingqian, Y., Donghong, L., Jianchu, C. & Peilong, S. (2013) Characterization of chemical composition of bee pollen in China. Journal of Agricultural and Food Chemistry, 61, 708-718.

Accepted 12 June 2020

Associate Editor: Sophie Evison