Ovary activation correlates with the reproductive potential of honeybee (*Apis mellifera*) workers if they are in a foreign colony

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Abstract

In eusocial Hymenoptera, the laying of male-determined eggs by workers in an unrelated colony can be a powerful strategy for increasing direct fitness benefits. A recent study showed that honeybee rebels, which are workers that develop under queenless conditions and have high reproductive potential, drift to foreign colonies, with a preference for hopelessly queenless colonies, and act as reproductive parasites. In our experiment, by introducing 5-day-old rebels and normal workers into foreign queenright colonies, we discovered that only rebels developed a reproductive phenotype by activating their ovaries. In a similar experiment with 1-day-old workers, neither rebels nor normal workers displayed such a tendency. We suggest that workers’ reproductive potential could thus be a key parameter activating the ovaries in not only drifting workers but also workers acting as reproductive parasites. Our results also support the hypothesis that the colony recognition cue probably has an environmental, rather than a genetic, origin.

Keywords Intraspecific reproductive parasites · Rebel workers · Ovary activation · *Apis mellifera*

Introduction

One of the most prominent features of hymenopteran insect societies is the reproductive division of labour. Typically, queens monopolize reproduction, while workers are facultatively sterile and refrain from reproducing in the presence of the queen (Wilson 1971; Bourke 1988). However, honeybee (*Apis mellifera*) colonies are vulnerable to exploitation by cheater workers that lay eggs to be reared either in their own colony or in the colony of conspecifics. This latter phenomenon is called intraspecific reproductive parasitism and is fairly common in western honeybees (Neumann and Moritz 2002; Moritz and Neumann 2004; Pirk et al. 2007; Beekman and Oldroyd 2008; Kuszewska et al. 2018) and in other *Apis* species (Paar et al. 2002; Nanork et al. 2005, 2007; Chapman et al. 2009).

In honeybees, workers drift between colonies, and in some apiaries, up to 40% of the workers in a colony are from foreign nests (Pfeiffer and Crailsheim 1998). Initially, failures in orientation caused by short distances between colonies in the apiary were thought to be a major factor explaining the presence of nonnatal workers in the colony (Jay 1965; Pfeiffer and Crailsheim 1998). Drifting individuals were expected to be sterile because the pheromones of the queen efficiently inhibit the development of ovaries in workers (Winston 1987). However, another study has shown that workers drifting to queenless colonies have disproportionately higher reproductive success than native individuals (Chapman et al. 2010). The absence of a queen and her pheromones is the most common factor stimulating ovary activation in workers (Winston 1987) and switches off the normal policing mechanisms, which is the removal of worker-laid eggs (Woyciechowski and Łomnicki 1987; Ratnieks and Visscher 1989; Miller and Ratnieks 2001). Consequently, workers that drift to a queenless colony may have a better chance of reproducing than workers that drift to a queenright colony. However, a study by Chapman et al. (2010) showed that drifting workers had no preference for queenless or queenright colonies, suggesting that this phenomenon is the result of mistakes in orientation. Different study results have shown that workers tend to migrate to foreign colonies, with a preference for hopelessly queenless colonies (Yagound et al. 2017). Similar results were obtained by Kuszewska et al. (2018), who studied the intraspecific reproductive parasitism of rebel workers in honeybees.
Rebel workers are a recently discovered subcaste of honeybee workers characterized by high reproductive potential (Woyciechowski and Kuszewska 2012). In comparison to typical workers, the rebels are more queen-like. They have significantly more ovarioles in their ovaries and larger mandibular and Dufour’s glands, but their hypopharyngeal glands, which synthesize and store brood food (Huang and Otis 1989), are underdeveloped (Woyciechowski and Kuszewska 2012; Kuszewska and Woyciechowski 2015). Additionally, studies of rebels have shown that they lay their own male-determined eggs more often than normal workers, even if they remain in a colony with a queen (Kuszewska et al. 2018). The proximate factors of rebel subcaste development are environmental conditions experienced during the larval stage. Rebels develop in the absence of a queen or, more precisely, in the absence of the queen’s mandibular gland pheromones (Woyciechowski and Kuszewska 2012; Woyciechowski et al. 2017). Naturally, such individuals occur after swarming, which is the only natural means of colony multiplication (Winston 1987). The ultimate factor explaining the rebel strategy is based on the assumption of inclusive fitness theory (Hamilton 1964) and results from a dramatic decrease in relatedness between the old queen’s workers and the new sister queen’s offspring after swarming (Woyciechowski and Kuszewska 2012). The high reproductive potential of rebel workers and their tendency to activate ovaries, even in queenright colonies, influences their behaviour and increases the possibility that they will drift to foreign colonies to lay unfertilized eggs. Indeed, rebels are more likely than normal workers not only to drift to foreign colonies but also to prefer colonies that are queenless (Kuszewska et al. 2018). These findings shed new light on intraspecific reproductive parasitism in honeybees; however, the mechanism underlying this strategy is not fully understood.

Obviously, if workers deliberately emigrate to lay eggs in a foreign colony, there must be some signal or cue that varies between colonies that can be perceived by workers. Evidence suggests that honeybees use both the blend of hydrocarbons present in their cuticle (Page et al. 1991; Châline et al. 2005; Zweden and d’Ettorre 2010) and fatty acids (Breed 1998; Breed et al. 2004) for chemical recognition of nest membership. The recognition cues are colony-specific and vary in the relative amount of individual components between colonies (Zweden and d’Ettorre 2010; Vernier et al. 2019). It has been hypothesized that transfer of the recognition cues between colony members occurs via exposure to the comb wax (Breed 1998; Couvillon et al. 2007; Zweden and d’Ettorre 2010). A recent study showed that differences in the chemical profiles of individuals from different honeybee colonies are a result of gut microbiota differences between colonies (Vernier et al. 2020).

Here, we tested whether rebels experimentally introduced into foreign queenright colonies perceive that they are not in a native colony and respond by activating their ovaries more often than normal workers who accidentally enter a foreign colony and behave in a similar way to natal nestmates (Smith and Loope 2016). We expected that rebels with high reproductive potential and a tendency towards intraspecific reproductive parasitism would be more prone to activate their ovaries after entering foreign colonies. The following predictions were tested: (1) foreign rebels have more developed ovaries than native rebels; (2) foreign normal workers do not differ in ovary activation from native normal workers; and (3) foreign and native rebels have more developed ovaries than foreign and native normal workers. Thus, the expected gradient of ovary development was foreign rebels > native rebels > foreign normal workers = native normal workers. We also examined whether workers learn colony recognition cues at the larval stage or as adult bees. Two hypotheses regarding the time of learning the colony recognition cues by workers were tested: (1) condition corresponds to larval-stage-experienced bees, while (2) condition corresponds to adult-stage-experienced bees.

Material and methods

The research was conducted in May and June 2018 in an experimental apiary at the Institute of Environmental Sciences of Jagiellonian University. Four queenright honeybee colonies with naturally mated queens were studied, each consisting of 20,000–40,000 workers. Each colony was treated in the same way as previously described (e.g. Woyciechowski and Kuszewska 2012; Kuszewska and Woyciechowski 2015), and the experiment was conducted over 2 successive days, with two colonies per day. Initially, rebel and normal workers were reared in all colonies (Fig. 1). For this purpose, the queen was confined to two experimental frames for 24 h to produce eggs of similar ages (day 0). Next, the colony was divided into queenright and queenless subunits, with each subunit containing one experimental frame (day 3). The subunits were reunited when the worker cells on the experimental frames were sealed (day 13), and beginning at that time, the experimental broods experienced the same conditions during their prepupal and pupal stages. Before the first adult workers emerged (day 21), the experimental frames were placed in an incubator in the laboratory (34 °C, 90% RH). 200 freshly emerged rebels (developing in a queenless subunit) and 200 normal workers (developing in a queenright subunit) from each experimental frame were marked on the thorax with a spot of paint (Marabu—Brilliant Painter). Afterwards, both rebel and normal workers were assigned to the following experimental groups: (1) introduced to the native colony on the 1st day of adult life; (2)
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| Day of experiment | Treatments |
|-------------------|------------|
| 0                 | Isolation of queen on two experimental frames in each of four experimental colonies of the experiment |
|                   | Colony 1  | Colony 2  | Colony 3  | Colony 4  |
|                   | Q        | R        | Q        | R        |
| 3                 | Division of each colony into two subunits: queenright (QR; rearing normal workers) and queenless (QL; rearing rebel workers) |
|                   | N        | R        | N        | R        |
|                   | Q        | R        | Q        | R        |
| 13                | Sealing of the cells with larvae on the experimental frames. Reunion of each of the subunits, thereafter both normal and rebel workers are placed again in QR colonies |
|                   | N        | R        | N        | R        |
|                   | Q        | R        | Q        | R        |
| 21                | Moving the experimental frames from each colony to an incubator (36 °C) and marking of emerging normal and rebel workers |
|                   | N        | R        | N        | R        |
|                   | Q        | R        | Q        | R        |
| 21                | Assignment the labelled normal and rebel workers to the following experimental groups: |
|                   | 1 – introduced to the native colony on the 1st day of adult life |
|                   | N        | R        | N        | R        |
|                   | Q        | R        | Q        | R        |
| 21                | 2 – introduced to the foreign colony on the 1st day of adult life |
|                   | N        | R        | N        | R        |
|                   | Q        | R        | Q        | R        |
| 21                | 3 – introduced to the native colony on the 1st day of adult life and then transferred to the foreign colony on the 5th day of adult life |
|                   | N        | R        | N        | R        |
|                   | Q        | R        | Q        | R        |
| 36                | Killing and dissecting normal and rebel workers from all experimental groups to assess their anatomical parameters |

Fig. 1 Timeline of the experiment, showing the manipulations on successive days
introduced to the foreign colony on the 1st day of adult life; and (3) introduced to the native colony on the 1st day of adult life and then transferred to the foreign colony on the 5th day of adult life (Fig. 1). We chose 5-day-old workers for transfer to the foreign colony, as during this time, workers have several days to learn colony recognition cues, but they have not yet begun foraging (Winston 1987). The types of workers (rebels vs. normal workers) and the experimental groups (1–3) were marked with different colours.

When the marked workers were 15 days old, 30 rebels and 30 normal workers were captured from each experimental group (1–3) to assess their number of ovarioles and ovary activation. We dissected the ovaries of the frozen workers and examined them under a stereomicroscope. We counted the total number of ovarioles in both ovaries (sum) and evaluated ovary activation by measuring maximal diameters of the most developed ovariole from each of the ovaries as described by Nakaoa et al. (2008), according to whom ovariole diameter reflects ovarian activity. All organs were stained with Giemsa reagent for approximately 10 s before being measured.

To compare the number of ovarioles between the rebel and normal workers, a mixed-model three-way ANOVA was used with the type of worker (rebel vs. normal) and the experimental group (1–3) as fixed effects and the colony of origin as a random effect. Statistically significant ANOVA results were followed by multiple comparisons using a post hoc Tukey honest significant difference (HSD) test, with \( p < 0.05 \) considered significant. We used a nonparametric Kruskal–Wallis test to compare the sizes of ovarioles from the two groups of workers and the different experimental groups (1–3) in each of the four colonies. All analyses were performed in Statistica 13.3 (StatSoft, Poland).

## Results

The results showed that rebels, which developed in queenless conditions at the larval stage, had more ovarioles in their ovaries than normal workers (three-way ANOVA, type of worker: \( df = 1, F = 261.808, p < 0.0005 \); Fig. 2a), which is consistent with the findings of previous studies (Woyciechowski and Kuszewska 2012; Kuszewska and Woyciechowski 2015; Kuszewska et al. 2018). Neither the colony of origin nor the experimental group (1–3) had a significant effect on the number of ovarioles in workers (three-way ANOVA; colony of origin: \( df = 3, F = 1.916, p = 0.2953 \); experimental group: \( df = 2, F = 0.942, p = 0.4404 \); Fig. 2a).

There were no differences in ovary activation between workers with different colonies of origin (Kruskal–Wallis test: \( p = 0.1490 \)). Ovarian development depended on the type of worker: rebels had more activated ovarioles than normal workers (Kruskal–Wallis \( p < 0.0001 \); Fig. 2b). Ovary activation did not differ between normal workers from groups introduced to the native and foreign colonies on the 1st day of adult life (\( p = 0.6056 \)), and workers from these groups did not differ from those in the group introduced to the foreign colony on the 5th day of adult life (\( p = 0.1888, p = 0.1326 \), respectively; Fig. 2b). There were no significant differences in ovary development between rebels from groups introduced to the native and foreign colonies on the 1st day of adult life (\( p = 0.7920 \), but the rebel workers from the group introduced to the foreign colony on the 5th day of adult life had significantly more developed ovarioles than rebels from the groups introduced to the native (\( p < 0.0001 \) and foreign colonies (\( p = 0.0058 \)) on the 1st day of their adult life.

## Discussion

In line with our predictions, workers with high reproductive potential, that is, rebels, responded to finding themselves in a foreign queenright colony by activating their ovaries if they had been introduced there on the 5th day of their adult life. In contrast, foreign normal workers, independent of the age at which they were moved, had the same low degree of ovary activation as their natal counterparts regardless of their age. These results suggest that the reproductive potential of workers affects their propensity to activate their ovaries in response to drifting behaviour.

Laying eggs in an unrelated colony can be a powerful strategy for increasing the direct fitness of workers (Beekman and Oldroyd 2008). Recent studies have shown that honeybee workers drift to foreign colonies, with a preference for hopelessly queenless colonies (Yagound et al. 2017; Kuszewska et al. 2018), which implies that acting as reproductive parasites is another option for workers’ life strategies. Our study shows that even if rebel workers are experimentally introduced into foreign queenright colonies, they activate their ovaries and presumably invest in personal reproduction in host colonies, while normal workers in the same situation do not. This outcome supports the findings of Kuszewska et al. (2018) that workers with high reproductive potential are more predisposed to function as reproductive parasites. Such behaviour is reasonable because rebels use the resources of unrelated colonies to raise their own male offspring, while their relatives do not bear the cost of caring for their sons.

Unlike rebels, normal workers did not activate their ovaries after being experimentally introduced into foreign queenright colonies. This finding is consistent with that of Smith and Loope (2016), who reported that normal workers entering foreign colonies due to orientation errors continued to refrain from personal reproduction, suggesting that they are caught in an evolutionary trap in which formerly reliable
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In turn, Yagound et al. (2017) showed that the few normal workers that drifted from queenright colonies were significantly more likely to have activated ovaries than their nondrifting natal nestmates. However, in their experiment, unlike ours and that of Smith and Loope (2016), workers followed deliberate reproductive drifting rather than nonreproductive accidental drifting, which are quite different phenomena (Nonacs 2017). Based on our results, the ability of workers to activate their ovaries in foreign colonies lies in their ability to react to the social context. It seems that high reproductive potential constitutes a key parameter promoting the acquisition of a parasitic phenotype in foreign queenright colonies. Similar outcomes were reported in workers in other eusocial insects. Bumblebee workers with active ovaries introduced into a foreign nest maintained their fertility, despite more frequent direct contacts with the queen and the brood than control workers. Conversely, infertile workers failed to develop a reproductive phenotype in a foreign nest (Yagound et al. 2012; Blacher et al. 2013). The study of intraspecific worker parasitism in the common wasp *Vespula vulgaris* showed that in foreign queenright colonies, drifters activated their ovaries at a rate that was ca. five times higher than the natal workers (Oliveira et al. 2016).

Another likely explanation for ovary activation by rebels in foreign queenright colonies is their greater learning ability. Learning allows individuals to adjust their behaviour to changing environmental conditions (Dukas 2008), which can have a considerable positive impact on individual fitness components. According to a recent study (Kuszewska and Rojek 2021), rebel workers learn faster than normal workers. As a result, the rebel workers in our...
study learned colony recognition cues before they were experimentally introduced on the 5th day of adult life to the foreign colony, which enabled them to detect changes in the colony and adjust their ovarian development, while normal workers presumably did not. However, we cannot exclude the alternative scenario in which normal workers detect themselves as being in a foreign queenright colony as rebel workers do, but they do not activate their ovaries due to lower physiological abilities or a longer time needed for activation. Some studies have shown that ovary activation is related to the number of ovarioles (Makert et al. 2006), and bees with more ovarioles are most likely to activate their ovaries (Amdam et al. 2006; Linksvayer et al. 2009; Wang et al. 2010; Traynor et al. 2014).

Interestingly, none of the groups of workers activated their ovaries after introduction into foreign colonies on the 1st day of adult life, which suggests that these workers did not recognize the change in the colony. Although some studies have shown that the recognition cue is acquired prior to emergence as an adult (Breed 1983), the experiments described in this paper support the hypothesis that the cue has an environmental, rather than a genetic, origin (Downs and Ratnieks 1999; D’Ettorre et al. 2006; Vernier et al. 2020). As the workers in our study were introduced to foreign colonies directly after emerging in the incubator, they probably recognized this colony as their own. Only after the 5th day of being in a given colony were they able to recognize whether the colony had changed. There might be no adaptation to emerge in foreign nests.

In conclusion, in our experiment, the rebels responded to finding themselves in a foreign queenright colony by activating their ovaries, while the normal workers did not employ such a strategy. This finding suggests that the particular reproductive potential of workers may be a prerequisite for ovary activation in drifting behaviour and that rebels with high reproductive potential are more predisposed than normal workers to function as reproductive parasites in conspecific colonies.

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Data availability The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.

Declarations

Conflict of interest The authors declare that they do not have any conflicts of interest.

Code availability Not applicable.

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