Emergency queen rearing in honeybee colonies with brood of known age

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Abstract – In four honeybee colonies, queens were isolated on empty combs for 8 consecutive days, so that in every colony there were 8 combs containing brood of known age. Afterwards, the colonies were dequeened and the process of emergency queen rearing was observed. The average interval from egg laying to queen cell capping was 8.8 days and ranged from 7 to 12 days. The average interval from queen cell capping to queen emergence was 7.2 days and ranged from 5 to 8 days. The whole development time from egg laying to queen emergence was 15.7 days, ranging from 14 to 18 days. The age of brood at the moment of dequeening positively correlated with both the time of capping and the total queen development time. The average age of brood (at time of dequeening) around which queen cells were built was 3.0 days. However, higher proportions of queen cells with younger larvae were destroyed; in effect, the age of brood at dequeening from which queens emerged was 3.4 days.

Apis mellifera / honeybee / queen rearing / development time

1. INTRODUCTION

Honeybee (Apis mellifera L.) workers in queenless colonies are able to rear queens from larvae primarily destined to be workers (Winston, 1979; Fell and Morse, 1984). Larvae for queen rearing often are available in a range of ages. The reproductive quality (as defined by Tarpy et al., 2000) of queens reared from younger larvae can be higher (Eckert, 1937; Boch and Jamieson, 1960; Weaver, 1957; Woyke, 1971; Tarpy et al., 2000). On the other hand, rearing queens from older larvae can shorten the queenlessness period (Tarpy et al., 2000). Hatch et al. (1999) demonstrated that honeybee workers rear emergency queens from brood that at the time of dequeening are 1 to 5 days old. However, their experimental setup did not allow the age of brood in queen cells to be determined directly. Instead they estimated it based on the time of queen cell capping, assuming that queen cells are capped 8 days after egg laying (Winston, 1987). This type of estimation has often been used in studies of emergency queen rearing (Winston, 1979; Fell and Morse, 1984; Hatch et al., 1999; Schneider and DeGrandi-Hoffman, 2002). The accuracy of this estimation has never been determined even though it is known that the time of queen cell capping varies considerably. Jay (1963) reviewed the literature concerning the duration of queen development times, reporting that queen cells are capped between 7 and 9 days after egg laying and that the whole development time varies from 15 to 17 days. Genotype is a major factor affecting queen development time. There are marked differences between subspecies (Fletcher, 1978; Winston, 1979; DeGrandi-Hoffman et al., 1998) and some variation between colonies of the same subspecies (Visscher, 1986; Tarpy and Fletcher, 1998; Hatch et al., 1999). Even within the same colony, temperature (DeGrandi-Hoffman et al., 1993) and nutrition (Visscher, 1986) affect queen

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development time. Other factors that are related to temperature and nutrition, such as the position of the queen cell in the nest (Visscher, 1986; Fell and Morse, 1984; DeGrandi-Hoffman et al., 1993; Hatch et al., 1999), affect development time as well.

Not all queen cells initiated by workers during an emergency queen rearing process successfully emerge. They can be destroyed both before and after capping (Allen, 1956; Caron and Greve, 1979; Schneider et al., 2001, 2002). The destruction of capped queen cells can be carried out by both workers and newly emerged queens (Allen, 1956; Fletcher, 1978; Caron and Greve, 1979; Gilley, 2001). Queens cut a small hole in the side wall of the queen cell and sometimes sting its occupant; workers cut a much bigger hole in the side wall and instead of stinging they remove the contents of the queen cell (Allen, 1956; Fletcher, 1978; Caron and Greve, 1979). Hatch et al. (1999) did not allow newly emerged queens to walk freely in the nest and destroy unemerged competitors in queen cells. Isolation of queen cells can affect the chances of queens being reared from brood of different ages, and in consequence, affect the reproductive quality of the only queen remaining in a colony when the process of emergency queen rearing is finished. Thus, to fully evaluate the factors influencing queen rearing, it is necessary to examine queen replacement under natural conditions in colonies containing brood of known age.

The purpose of this paper was to investigate emergency queen rearing in honeybee colonies. We had two main objectives. First, we determined the variability of the different stages of queen development and the accuracy of estimation of ages of brood in queen cells based on their time of capping. Second, we measured more accurately than previous studies the ages of brood from which emergency queens are produced. We examined the process of emergency queen rearing in colonies in which the age of young brood is known and newly emerged queens have access to unemerged queen cells.

2. MATERIALS AND METHODS

The experiment was carried out in June and July 2000, and used four colonies of honeybees considered to be Apis mellifera carnica. The colonies were of similar size, and each occupied two boxes containing 10 frames apiece. The frames were of the wielkopolski type locally used in Poland (width 360 mm, height 260 mm). To obtain eggs of known age, the queen from each colony was confined for 24 hours on an empty frame of comb in a cage made of wood and queen excluder. The caged queens were placed in their original colonies in the center of the bottom box. This procedure was repeated 8 times. Every day the queens were isolated on new frames of empty comb. At the end of the procedure all colonies were dequeened. At that time in the bottom box of each hive there were 8 combs containing brood ranging in age from 1-day old eggs to 5-day old larvae. The other two frames in the bottom box and the frames in the upper box contained brood older than 8 days, honey and pollen stores. The combs with brood of different ages were placed in a different order in each colony (Fig. 1). After removal of the queens, the colonies were examined every day at the same time of day until all queen cells were either emerged or destroyed. During inspections the position of every queen cell was marked on acetate sheets, one sheet for each side of each frame. If a regular round opening was found at the bottom of the queen cell it was categorized as emerged. If an opening was found in the side of the queen cell it was categorized as destroyed. We call the interval from egg laying to queen cell capping the precapping period, and the interval from queen cell capping to queen emergence the postcapping period.

In the statistical analysis we combined the data from all four colonies because we were more interested in variation within the population than in
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differences between colonies. The times of queen cell initiation, queen cell capping and queen emergence were analyzed in relation to both the time of dequeening and the time of egg laying. Associations between two variables (e.g., length of precapping period and distance from the bottom of the hive) were tested with Spearman's rank correlation. Differences between all continuous variables (e.g., length of precapping period) and discontinuous variables (e.g., number of queen cells) were analyzed using nonparametric tests: the Mann-Whitney test in the case of two groups and the Kruskal-Wallis test when there were more than two groups. The proportions of queen cells destroyed were analyzed using the G-test of independence (Sokal and Rohlf, 1995). The G-test cannot be calculated for frequency equal to zero, so the numbers of emerged and destroyed queen cells were combined across ages 6 to 8 days. All average values are reported as ± 1 SD.

3. RESULTS

3.1. General information

The numbers of initiated, capped and emerged queen cells per colony were 32.7 ± 4.79, 27.0 ± 1.83 and 13.0 ± 1.63 (N = 4), respectively. The numbers of queen cells on central and marginal frames were 19.0 ± 2.45 and 13.7 ± 2.63 (N = 4), respectively. Those values differ significantly (Mann-Whitney test: U = 0.5, N1 = N2 = 4, P = 0.028). The frame area covered by brood was 1.55 ± 0.55 dm² (N = 32), ranging from 0.20 to 2.76 dm². There was no significant correlation between the frame area covered by brood and the number of queen cells on the frame (Spearman’s rank correlation: rₛ = 0.208, N = 32, P = 0.254). The brood area did not differ between frames of different brood age (Kruskal-Wallis test: H = 5.18, df = 7, N = 32, P = 0.638), nor between central and marginal frames (Mann-Whitney test: U = 126, N₁ = N₂ = 16, P = 0.940).

3.2. Precapping period

The queen cell precapping period was 8.8 ± 1.30 days (N = 108) and ranged from 7 to 12 days (Fig. 2B). Queen cells from which queens emerged were capped at a slightly, but significantly earlier age (8.5 ± 1.08 days; N = 52) than those that were destroyed (9.1 ± 1.43 days; N = 56) (Mann-Whitney test: U = 1114, N₁ = 52, N₂ = 56, P = 0.020; Fig. 2B). The precapping period of queen cells from which queens emerged positively correlated with the age of brood at the time of dequeening (Spearman’s rank correlation: rₛ = 0.555, N = 52, P < 0.001; Fig. 3A) and with the distance of the queen cell from the central frame (Spearman’s rank correlation: rₛ = 0.317, N = 52, P = 0.022). The precapping period negatively correlated with the distance of the queen cell from the bottom of the hive (Spearman’s rank correlation: rₛ = –0.304, N = 108, P = 0.001), but this relationship was not significant when only queen cells from which queens emerged were analyzed (Spearman’s rank correlation: rₛ = –0.141, N = 52, P = 0.318). The precapping period negatively correlated with the postcapping period

Figure 2. Distribution of ages of brood around which queen cells were built (A), distribution of ages of brood at time of queen cell capping (B), distribution of ages of brood at time of queen cell destruction (C), and distribution of whole development times from egg laying to queen emergence (D) during emergency queen rearing. Bars represent averages ± SD across four colonies.
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The precapping period of queens reared from brood, which was younger at the time of dequeening, tended to be shorter than the precapping period of queens reared from older brood. The queen cells in the center of the nest tended to be capped after a shorter period of time than the queen cells at the nest periphery.

3.3. Postcapping period

The postcapping period was 7.2 ± 0.78 days (N = 52) and ranged from 5 to 8 days (Fig. 3B). The postcapping period negatively correlated with the distance of the queen cell from the central frame (Spearman’s rank correlation: \( r_s = -0.311, N = 52, P = 0.025 \)). There was no significant relationship between the postcapping period and the age of brood at the time of dequeening (Spearman’s rank correlation: \( r_s = -0.432, N = 52, P = 0.001 \)). The postcapping period of queens reared in the center of the nest tended to be longer than the postcapping period of queens reared at the nest periphery but was not affected by the age of brood at the time of dequeening.

3.4. Total development time

Queen emergence occurred between 14 and 18 days after the egg was laid, with an average of 15.7 ± 0.88 days (N = 52; Fig. 2D). The total development time positively correlated with the age of brood at the time of dequeening (Spearman’s rank correlation: \( r_s = 0.561, N = 52, P < 0.001 \); Fig. 3C). There was no significant relationship between the total development time and the distance of the queen cell from the central frame (Spearman’s rank correlation: \( r_s = 0.118, N = 52, P = 0.406 \)) and the distance of the queen cell from the bottom of the hive (Spearman’s rank correlation: \( r_s = -0.017, N = 52, P = 0.904 \)). The total development time of queens reared from brood that was younger at the time of dequeening tended to be shorter than the total development time of queens reared from older brood, but did not depend on the position of the queen cells in the nest.

3.5. Initiation and destruction of queen cells

The queen cells were initiated around brood aged between 3 and 11 days; the average age of brood used to initiate queen cells was 5.9 ± 1.90 days (N = 131; Fig. 2A). Eggs were never observed inside queen cells. Queen cells from which queens emerged and those destroyed either before or after capping did not differ in the age of brood from which they were constructed (Mann-Whitney test: \( U = 1732, N_1 = 52, N_2 = 79, P = 0.123 \); Fig. 2A). Most of the queen cells (60.3%) were destroyed, 17.6% of them before capping and 42.7% after capping (Fig. 2C). The queen cells were destroyed between the 5th and 18th days of brood development, and the average age of brood at the time of queen cell destruction was 13.0 ± 3.48 days (N = 79; Fig. 2C). A smaller
3.6. Use of brood of different ages

The age (at time of dequeening) of brood around which queen cells were built and from which queens emerged was $3.0 \pm 1.40$ (N = 131) and $3.4 \pm 1.27$ (N = 52) days, respectively (Fig. 4). Queen cells from which queens emerged were initiated using brood that was older at the time of dequeening ($3.4 \pm 1.27$ days; N = 52) than the brood of those destroyed either before or after capping ($2.7 \pm 1.40$ days; N = 79) (Mann-Whitney test: U = 1327, N₁ = 52, N₂ = 79, $P < 0.001$; Fig. 4). The age of brood at the time of dequeening significantly affected the number of queen cells constructed using the brood (Kruskal-Wallis test: H = 21.2, df = 7, N = 32, $P < 0.004$; Fig. 4). The greatest number of queen cells were constructed over brood 3 days old at the time of dequeening (Fig. 4). The age of brood at the time of dequeening also affected the proportion of queen cells destroyed (G-test of independence: G = 17.0, df = 5, $P = 0.005$; Fig. 4). The smallest proportion of queen cells were destroyed on frames with brood 4 days old at the time of dequeening (Fig. 4).

3.7. Emergency queen rearing in relation to time of dequeening

Queen cells were initiated between the 1st and 9th days after dequeening (Fig. 5A). In two of the colonies the first queen cells were observed the day after dequeening, and in the other two colonies the second day after dequeening. Queen cells from which queens emerged were initiated significantly earlier than those destroyed either before or after capping (Mann-Whitney test: U = 1372, N₁ = 52, N₂ = 79, $P = 0.001$; Fig. 5A). The distribution of times at which queen cells were initiated was bimodal, with the first mode on the 2nd–3rd day after dequeening and the second mode on the 9th day after dequeening (Fig. 5A). The queen cells were capped between the 4th and 11th days after dequeening, and the peak of queen cell capping occurred the 6th–7th day after dequeening (Fig. 5B). The queen cells from which queens emerged were sealed significantly earlier than those destroyed after capping (Mann-Whitney test: U = 752, N₁ = 52, N₂ = 56, $P < 0.001$; Fig. 5B). Queens emerged
from the queen cells between the 11th and 16th days after dequeening, and the peak of queen emergence occurred the 13th day after dequeening (Fig. 5D). Queens reared from brood older at the time of dequeening tend to emerge earlier than queens reared from younger brood (Spearman’s rank correlation: \( r_s = -0.758, N = 52, P < 0.001 \)). Destruction of queen cells was observed between the 3rd and 16th days after dequeening (Fig. 5C). The distribution of times at which queen cells were destroyed was bimodal, with the first mode on the 4th–6th day after dequeening and the second on the 12th day after dequeening (Fig. 5C).

4. DISCUSSION

Our data show that both the precapping period and the whole development time of emergency queens increase with the age of brood from which the queens were reared (Fig. 3). Thus, estimations of brood age in emergency queen cells based on time of capping (Winston, 1979; Fell and Morse, 1984; Hatch et al., 1999; Schneider and DeGrandi-Hoffman, 2002) cannot be very accurate. Another source of inaccuracy of estimations is the correlation of the precapping period with the position of the queen cell in the nest (Visscher, 1986; Fell and Morse, 1984; DeGrandi-Hoffman et al., 1993; Hatch et al., 1999). Both the distance from the central frame and the distance from the bottom of the nest affected the length of the precapping period in this study. The differences in the length of the precapping period of brood in queen cells positioned in different parts of the nest are compensated by the length of the postcapping period. In consequence the whole development time does not depend on the position of the queen cell in the nest. Bienefeld (1996) observed a similar negative correlation between the precapping and postcapping periods in honeybee workers.

Estimating age of brood in queen cells on the assumption that the precapping period is always 8 days often results in underestimations of brood age. In the majority of cases the underestimation is low compared to the error of one day accepted by most studies of emergency queen rearing (Hatch et al., 1999; Schneider and DeGrandi-Hoffman, 2002). For the average age of brood used to produce queens in this study (2.96 days; Fig. 4), the difference between the estimated and actual precapping period (calculated using linear regression coefficients; Fig. 3A) was 0.80 days. However, some of the queen cells were capped 12 days after egg laying (Fig. 2B). In those rare (3%) cases the error of estimation of the age of brood in queen cells can reach four days. During swarming, all queen cells are initiated with newly laid eggs, so brood age estimations based on the time of capping can then be much more accurate, although this needs to be verified experimentally. Other factor affecting the accuracy of the estimation will be the position of the queen cell in the nest. Our results suggest that the whole development time, instead of the time of capping, should be used for estimation because it does not depend on the position of the queen cell in the nest.

We studied emergency queen rearing under natural conditions and did not isolate the capped queen cells. This means that the time of emergence of queens was affected not only by the age of brood (at time of dequeening) and the position of queen cells in the nest but also by other factors that can influence queen emergence and which may differ between emergency queen rearing and swarming. Workers standing on queen cells often perform a vibration signal, which consists of dorsoventral vibration of their body (Allen, 1959; Painter-Kurt and Schneider, 1998). It has been suggested that the vibration signal affects timing of queen emergence (Fletcher, 1978; Bruinsma et al., 1981; but see Grooters, 1987; Schneider et al., 2001) and inhibits interactions between queens (Fletcher, 1978; Schneider, 1990, 1991). Moreover, emerged queens produce a series of pulsed sounds called piping which can delay the emergence of other queens (Grooters, 1987). Piping is more common during swarming than during emergency queen rearing (personal observations). A factor probably limited to swarming periods is imprisonment of young queens in queen cells (Fletcher, 1978; Bruinsma et al., 1981). The mentioned factors may influence the timing of queen emergence and thus may have affected the positive relationship between age of brood (at time of dequeening) and the total development time. Other experimental setups and particularly isolation of capped queen cells in an incubator can influence the length of different stages of queens development. This has to be taken into account when the data presented here are compared with results of other studies.
The average lengths of different stages of honeybee queen development from our study agree with those from other studies (Jay, 1963; Winston, 1987), but the variation was greater in our experiment (Fig. 2). The large variation can be explained partly by differences in measurement accuracy, which in our study was affected by the length of time between consecutive queen cell inspections (24 h) and also by the length of time provided for queens to lay eggs in a single frame of comb (24 h). It is difficult to obtain higher accuracy of brood age measurements in a full-size colony, because frequent inspections disturb it and can affect the results.

We showed that workers initiate queen cells using brood in a wide range of ages at dequeening, but the greatest number of queen cells were produced around brood that at the time of dequeening were 3 days old (Fig. 4). Hatch et al. (1999) reported a similar pattern of queen cell initiation, but the range of ages used to produce queen cells was narrower in their study. This discrepancy is probably due to inaccurate estimates of brood age, based on the time of queen cell capping. Even though newly emerged queens are able to destroy other queen cells, in our experiment a considerable number of queens emerged in each of the colonies. Either workers protected the queen cells from destruction (Gilley, 2001) or the early emerged queens could not find and destroy the remaining queen cells before the queens emerged from them. Because queen cells with younger brood are more often destroyed, the average age of brood (at time of dequeening) from which queens emerged was higher than the average age of brood (at time of dequeening) around which queen cells were built (Fig. 4). We did not observe queen cell destruction in progress; another experiment is needed to determine whether the queen cells with younger brood were destroyed by newly emerged queens or workers. This problem needs to be addressed so that the mechanisms by which the quality of emergency queens is controlled in honeybee colonies can be better understood.

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