Changes in chemical cues of *Melissococcus plutonius* infected honey bee larvae

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**Abstract**

European foulbrood (EFB), caused by *Melissococcus plutonius*, is a globally distributed bacterial brood disease affecting *Apis mellifera* larvae. There is some evidence, even if under debate, that spreading of the disease within the colony is prevented by worker bees performing hygienic behaviour, including detection and removal of infected larvae. Olfactory cues (brood pheromones, signature mixtures, diagnostic substances) emitted by infected individuals may play a central role for hygienic bees to initiate the disease-specific behaviour. However, the mechanisms of cue detection and brood removal, causing hygienic behaviour in EFB affected colonies, are poorly understood. Here, coupled gas chromatography-mass spectrometry (GC–MS) was used to detect disease-specific substances, changes in cuticular hydrocarbon (CHC) profiles, and brood ester pheromones (BEPs) of honey bee larvae artificially infected with *M. plutonius*. Although no diagnostic substances were found in significant quantities, discriminant analysis revealed specific differences in CHC and BEP profiles of infected and healthy larvae. β-Ocimene, a volatile brood pheromone related to starvation and hygienic behaviour, was present in all larvae with highest quantities in healthy young larvae; whereas oleic acid, a non-volatile necromone, was present only in old infected larvae. Furthermore, γ-octalactone (newly discovered in *A. mellifera* in this study) was detectable in trace amounts only in infected larvae. We propose that the deviation from the olfactory profile of healthy brood is supposed to trigger hygienic behaviour in worker bees. To confirm the relevance of change in the chemical bouquet (CHCs, BEPs, γ-octalactone, etc.), a field colony bioassay is needed, using healthy brood and hygienic bees to determine if bouquet changes elicit hygienic behaviour.

**Keywords** *Apis mellifera* · European foulbrood · Gas chromatography-mass spectrometry · Cuticular hydrocarbons · Brood ester pheromones · Hygienic behaviour

**Introduction**

In the wake of the evolution of eusociality, hygienic behaviour has evolved as an important trait to control transmission and spread of diseases and parasites within social insect colonies (Cremer et al. 2007). In performing hygienic behaviour, worker honey bees, at the stage between nursing and foraging, detect and remove diseased larvae or pupae, even before they show visible symptoms (Evans and Spivak 2010). The two-step mechanism of hygienic behaviour includes not only the detection and removal of target individuals from open brood cells (bacteria and fungi-infected brood), but also from capped cells (mostly mite infested brood). The latter mechanism has an additional step, the uncapping of infested cells (removal of the wax lid) to get access to the brood and its parasite. The underlying mechanisms are based on olfactory cues to discriminate infected from healthy brood and to initiate brood removal. Both
leads to substantial worldwide colony losses (Ellis and Munn 2005). Especially in England and Wales, as well as Switzerland, disease outbreaks have risen rapidly through the last decades (Roetschi et al. 2008; Wilkins et al. 2007; von Büren et al. 2019). Local variation in disease outbreaks has been observed, which might be due to a variability of strain virulence (Budge et al. 2014).

Bailey (1960) observed several colonies performing hygienic behaviour on 2- to 10-day-old infected honey bee larvae. Significantly more treated larvae were removed by worker bees before pupation than control larvae. In his study, healthy 1-day-old larvae were artificially infected with *M. plutonius* on brood combs and placed back inside healthy colonies. Moreover, Bailey assumed that in natural disease outbreaks larvae die before pupation and are ejected earlier compared to artificially infected larvae (Bailey 1960, 1963). His assumption might be based on the fact that naturally infected colonies show signs of malnutrition and may respond differently from artificially infected colonies.

This study focuses on odorant changes in EFB-infected larvae that may serve as trigger for brood removal by hygienic worker bees. Honey bee larvae were artificially infected (Lewkowski and Erler 2019) using a highly virulent *M. plutonius* strain isolated in Switzerland (Djukic et al. 2018; Grossar et al. 2020; Lewkowski and Erler 2019). Chemical extracts and more precisely CHC profiles, BEP composition, oleic acid and *E*-β-ocimene of in vitro reared larvae were characterized using coupled gas chromatography-mass spectrometry (GC–MS) to identify potential diagnostic substances and changes in the odorant bouquet related to the disease. The degree of the disease was characterized phenotypically by larval age and colour change. Healthy and diseased bees were further characterized for presence and quantity of *M. plutonius*. Finally, disease symptoms and olfactory signals were analysed for correlations that may show at which stage the disease might be detectable by the worker bees.

### Material and methods

#### Larvae rearing, bacteria cultivation and infection

*Apis mellifera* worker larvae were reared in vitro according to the protocol of Crailsheim et al. (2013). A honey bee queen was caged on an empty brood frame for 24 h. Three days after releasing the queen, freshly hatched worker larvae (<24 h) were grafted and transferred into 24-well plates on 5 µl day–1 food solution, receiving food for 6 days (Crailsheim et al. 2013). To limit the variance of the larval bouquet (CHC profiles) to differences mainly resulting from the infection, only larvae of a single queen have been used for this study.
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*Melissococcus plutonius* (strain 49.3, wild type isolated in Switzerland, characterized in Djukic et al. 2018; Grossar et al. 2020) was cultivated in liquid medium (Lewkowski and Erler 2019) for 3 days, under semi-anaerobic conditions (10% CO₂). For larvae infection, bacteria-inoculated medium was mixed with day-1 food solution (Crailsheim et al. 2013) in a 1:10 ratio. The bacterial solution was previously adjusted to OD₆₀₀nm = 0.3. Effective number of colony-forming units (CFU) was estimated by CFU counting using serial dilutions (1:10, 1:100, 1:1000; Lewkowski and Erler 2019). The non-infected control group received day-1 food solution mixed with sterile culture medium at the same ratio. 5 µl of the infective food mixture was fed to 1-day-old worker larvae right after grafting (final infective dose: ~7.2 × 10³ CFUs per larvae). Eight days post-grafting, all larvae were weighed and transferred into new 24-well plates for pupation (Lewkowski and Erler 2019). For survival analysis dead individuals (larvae, pupae, pre-adult bees) were counted daily until eclosion. Infection success and potential pathogen variance among samples (larval age, larvae grouped to different colour classes) was tested using *M. plutonius* specific qPCR with an internal *A. mellifera* reference gene (COI) for normalisation (Lewkowski and Erler 2019), and with the primer set of Budge et al. (2010) (*EFBFor*–TGTTGTAGAGGAATAAGGGGAA, *EFBRev2*–CGTGGGTTTCTGGTTAGA). Genomic DNA of *M. plutonius* strain 49.3 was used as positive control, as well as a non-template water sample for the negative control.

**Sampling and gas chromatography-mass spectrometry (GC–MS)**

Larvae of different age (day 5, 8, 9, 10 and 15 post-grafting and infection) and colour (white, yellow, brown, black, Fig. 1) were sampled daily, starting with day 5 post-grafting (Table S1). As for infected larvae, larval age and disease-resulting colour change are partially correlated (on average older larvae show darker colour), not every time point was sampled with every larval colour (Table S1). Each ‘infected larvae’ replicate (n = 1–3) included three pooled larvae of the same age and colour in one glass vial according to the colour scheme shown in Fig. 1. At every time point, sampling diseased larvae, three replicates (with three larvae per replicate) of ‘non-infected control larvae’ were sampled as well. On day 8, after transferring larvae for pupation, remaining food, not consumed by the larvae, was pooled (due to low volume) and collected separately for each treatment group (infected, n = 6, control, n = 5). Remaining food, that also included some larvae faeces, was sampled to measure odorants secreted by the pathogen (*M. plutonius*) or the honey bee larvae at the feeding stage. Larvae or food remains (100 µl, respectively, per food replicate) were transferred into columnar AR-glasses (60 × 16.75 mm), covered with 200 µl *n*-hexane (95%, AppliChem) and extracted under medium shaking conditions for 10 min at room temperature. Samples were stored at −80 °C for minimum 24 h to prevent odorant evaporation, to generate an extract in equilibrium and to ensure sampling of the liquid extract phase (Seidelmann and Rolke 2019). Thereafter, 100 µl hexane extract (cuticular or food extract) were transferred into short thread ND9 glass vials (Carl Roth, Germany) and stored again at -80 °C until GC–MS analysis. The reduced volume was used to prevent solvent contamination with food particles or larval tissue.

Samples were analysed by a Saturn 2100 GC–MS (CP 3900 GC, Saturn 2100 T ion trap MS, Saturn-MS Workstation V 6.20; Varian Inc., California) equipped with an 1177 split/splitless injector and a VF-5 ms capillary column (30 m × 0.25 mm × 0.25 µm). Helium was used as carrier gas throughout. Oven temperature was programmed according to target substances (see below). Sample volumes of 1 µl were injected in splitless mode. Mass spectra were acquired using standard 70 eV electron impact ionization mode, mass range of 40–400 m/z. Both larvae and food odorants were analysed to characterize CHC-compounds. Low-boiling volatiles as BEP were furthermore analysed in selected high-content

![Fig. 1 Larval colour change during manifestation of the typical symptoms of EFB disease, from pearly white, yellow, brown to greyish black. Age (days post-grafting) of the larvae at each sampling point are marked with arrows. Three individuals of the same colour, according to this colour scheme, and same age were used for one GC–MS sample](image-url)
larvae samples. Substances have been identified by reference substances (e.g., alkanes C₇-C₄₀, BEP; Sigma-Aldrich) or their established substance-specific mass spectra (NIST MS-Database, Rev. 2.0) and those reported in the literature (Salvy et al. 2001), in conjunction with Kovats-indices determined using the alkane standard. For the identification of γ-octalactone (also known as 5-butylidihydro-2(3H)-furanone; 4-hydroxy octanoic acid; octano-1,4-lactone; γ-octanoic lactone; CAS 104-50-7), a synthetic reference substance (Sigma-Aldrich) was used.

Cuticular hydrocarbon profiles: CHC-compounds were identified and analysed according to the checklist and data provided by Salvy et al. (2001). GC was programmed to a linear temperature gradient of 10 °C/min, from 80 to 320 °C, 4 min at 320 °C, constant flow rate of 0.7 ml/min, and injector temperature 250 °C. Mass fragments were detected from 6 to 30 min.

Brood ester pheromone (BEP) composition and diagnostic substance: BEP profiles of honey bee larvae typically include methyl and ethyl esters of palmitic, stearic, oleic, linoleic and linolenic acid (Le Conte et al. 1989, 1990). Oleic acid (C₁₈H₃₄O₂) and (E)-β-ocimene (C₁₀H₁₆) have been discovered to be related to larval food-begging, larval starvation and brood removal by worker honey bees (He et al. 2016; Maisonnasse et al. 2010; McAfee et al. 2017, 2018). Under normal colony conditions, starving honey bee larvae emit higher amounts of the volatile brood pheromone β-ocimene and thus modulate the behaviour of worker bees to visit cells more frequently (He et al. 2016). β-Ocimene is also a major compound of the odour bouquet released by 0–3-day-old larvae, whereas 5-day-old larvae produce more of the low-volatile brood ester pheromones (BEPs) and vice versa (Maisonnasse et al. 2010). To quantify these lower boiling substances, the temperature program started at 40 °C with a linear gradient of 10 °C/min to 300 °C, injector temperature 240 °C, flow rate 1 ml/min, detection window of 5–30 min. Methyl linoleate was not detectable in any sample, and ethyl linoleate was detectable in a few samples, however not in quantifiable amounts (Fig. S1). Both compounds were omitted from further analysis. Methyl palmitate was measurable only in 36% of the samples in reliable quantities and had to be removed from the final data set as well.

Data analysis

Larval mortality was compared using Kaplan–Meier survival analysis following log-rank test. Larval weight and bacterial loads (M. plutonius qPCR) were tested by Shapiro–Wilk test for normal distribution, which could not be confirmed. Mann–Whitney U tests and Kruskal–Wallis ANOVAs (following Bonferroni adjusted Dunn’s post hoc test) were used to test for significant weight differences (mean ± SD) between infected and control larvae, as well as for variance in M. plutonius bacterial loads among samples. All data were analysed using Statistica 8.0 (StatSoft) and R (version 3.4.2).

Cuticular hydrocarbon GC–MS data (peak area and corresponding retention time), generated by automatic integration, were manually verified with due regard to mass spectra and exact retention times and manually re-integrated in case of deviating integration, were manually verified with due regard to mass spectra and exact retention times and manually re-integrated in case of deviation. Some compounds occurred randomly in few samples only in very low quantities and have been excluded from further analysis. Relevant compounds were checked for randomly (not age- or treatment-specific) distributed zero values and were excluded to generate a reliable dataset. The final dataset consisted of 32 compounds for each of the 25 larval samples. Peak areas of these compounds were converted by quantile normalization (‘preprocessCore’ package, Bolstad 2016; R Core Team 2017) and centred log-ratio transformation (Seidelmann and Rolke 2019) to achieve normal distribution. The R ‘stats’ package (R Core Team 2017) was used for performing a Principle Component Analysis (PCA). A discriminant function analysis was run based on all principle components, with eigenvalues above 1, to compare and visualize changes in the composition of cuticular hydrocarbon profiles of M. plutonius-infected and control larvae. Comparing the food samples, many substances could not be detected reliably and were listed as ‘zeros’. Consequently, a quantile normalization was not possible and substance raw peak area data were used for subsequent analysis. Four substances of the larvae data set could not be detected in food samples and were eliminated from the final food data set. Raw data of honey bee larvae BEP compounds were checked, normalized, transformed and analysed (PCA and discriminant function analysis) as described for the cuticular hydrocarbons.

Characteristic bouquet compounds were normalized to the pentacosane and heptacosane peak area (acting as internal standards), the two (n-C₂₅, n-C₂₇) major compounds of the honey bee larvae profile (Salvy et al. 2001). Normalized values were used to test for significant differences between age groups and/or larvae infected with M. plutonius vs. healthy, and to test for correlations between compound quantities, bacterial loads or age groups, respectively. Non-parametric tests were applied as data sets were not normally distributed.

Results

Melissococcus plutonius infection success was shown by treatment-dependent weight differences and larval survival. Infected individuals had a lower weight (95 ± 31 mg, n = 141) compared to controls (141 ± 20 mg, n = 131) (MWU test: U = 2142.5, p < 0.0001). Larval mortality at day 21 post-grafting was higher for infected individuals (76.5%,
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Infectious load (Spearman rank correlation, infected samples, γ-octalactone quantities increased with monoenes and dienes) (Table 1). Identified hydrocarbons saturated, branched and methyl-alkanes as well as alkenes compounds of the larval cuticular hydrocarbon bouquet were identified. The brood pheromone β-ocimene was present in all larvae showing much higher values (Cq median = 18.15) than day 8 larvae (Cq median = 19.38, which means a 3.73-times higher load for day 5) (z = 2.86, p = 0.042). All other time points did not differ from each other in M. plutonius loads. There was no significant interaction of larval colour (Fig. 1) and M. plutonius loads (Kruskal–Wallis ANOVA: H = 9.40, df = 4, n = 39, p = 0.002) (Table S2). Among infected samples, γ-octalactone quantities increased with increasing infection loads (Spearman rank correlation, r = 0.64, p < 0.01), but not with larval colour change (Spearman rank correlation, r = 0.06).

GC–MS data of larval extracts revealed that the main compounds of the larval cuticular hydrocarbon bouquet were saturated, branched and methyl-alkanes as well as alkenes (monoenes and dienes) (Table 1). Identified hydrocarbons ranged from chain lengths C_{17}–C_{31}. Cuticular hydrocarbon profiles of both treatment groups showed no differences in the identity of substances detected (Fig. 2). The PCA reduced the 32 components (peak no. 2–42) detectable in all control and infected individuals to seven factors explaining 85.36% of the total variance. The discriminant analysis separated larvae not only by age (day 5 vs. ≥ 8 post-grafting; Factor 1: 34.5% of total variance, eigenvalue: 11.03), but also by treatment (infected vs. control; Factor 2: 15.4% of total variance, eigenvalue: 4.94). The two factors represent about 50% of the total variability of the dataset. Few infected individuals cluster together with non-infected controls (Fig. 3). Larvae may survive the infection as shown by 23.5% larvae surviving until day 21 post-grafting (Fig. S2). One larva of the samples of day 8 yellow/brown and day 10 yellow/brown showed to be non-infected or had very low M. plutonius loads. With showing Cq-values of approx. 30 (see above) we can only speculate that the larvae were infected (without corresponding disease symptoms), or the bacteria proliferated at very low frequency, or these values came from feeding infectious food and the bacteria did not replicate. These single larvae may exude an odour bouquet like healthy larvae, grouping the whole sample to this specific cluster.

The main compounds contributing to the positive part of the first dimension, discriminating honey bee larvae by age, are long-chained linear alkanes (C_{27}–C_{29}, Table 1, Fig. S3). Whereas shorter (methyl-branched) alkanes represent the negative part of the first dimension (Table 1, Fig. 3, Fig. S3). Two CHC compounds strongly decreased with increasing larval age (Spearman rank correlation, C_{22}-docosane: r = −0.72, C_{25}-pentacosane: r = −0.93, both p < 0.001). The positive and negative part of the second dimension, discriminating the samples by treatment, was almost exclusively driven by odd (methyl-branched) alkanes and alkenes (C_{19}–C_{31}) (Table 1, Fig. 3, Fig. S3). For the food samples, the PCA reduced the 39 components to four factors explaining 93.41% of the total variance. The discriminant analysis separated food samples by treatment, with 56.8% (Factor 1, eigenvalue: 22.15) of the total variance (Fig. S4), albeit a high variance among non-infected control food samples can be observed.

The PCA for the seven BEP compounds reduced all components to three factors explaining 82.22% of the total variance. The discriminant analysis revealed one narrow group of old infected larvae (≥ day 8), surrounded by healthy old (≥ day 8) and healthy and infected young (day 5) larvae (Fig. 4). The first two factors explain 66.6% of the total variance (Factor 1, eigenvalue: 3.28; Factor 2, eigenvalue: 1.38). The clustering of the old infected larvae was mainly driven by ethyl oleate, ethyl palmitate and ethyl stearate (Fig. S5, Table S2). Comparing the different age groups showed that methyl oleate and methyl linolenate were much higher in younger (day 5) than in older (≥ day 8) larvae (MWU test: U_{methyl oleate} = 22.5, U_{methyl linolenate} = 23.5, n_{each} = 25, p < 0.05), whereas methyl stearate and ethyl stearate were much higher in older (≥ day 8) than in young larvae (day 5) (MWU test: U_{methyl stearate} = 25.5, U_{ethyl stearate} = 6, n_{each} = 25, p < 0.05) (Table S2).

The brood pheromone β-ocimene was present in all larvae samples, with younger larvae (day 5) showing much higher values than older (all larvae ≥ day 8) (MWU test: U = 6, p = 0.0013) (Table S2). Infection with *M. plutonius* decreased β-ocimene compared to healthy controls (MWU test: U = 30, p = 0.999).
$p = 0.019$ (Table S2), with no correlation between β-ocimene and *M. plutonius* loads for infected larvae (Spearman rank correlation, $r = 0.22$). Oleic acid was detectable only in larvae older than day 5 and infected with *M. plutonius* (Table S2), however without any significant differences between age groups (day 8–15, Kruskal–Wallis ANOVA: $H = 8.56$, df = 4, $n = 16$, $p = 0.07$) or any correlation with *M. plutonius* loads (Spearman rank correlation, $r = 0.12$).

### Discussion

Artificial infection of in vitro reared honey bee larvae generated phenotypes that were highly suitable to study differences in chemical bouquet composition of *A. mellifera* larvae. The following parameter (survival, colour, weight) made it possible to describe and group larval

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**Table 1** Main cuticular hydrocarbon profile compounds identified by GC–MS with peak number, compound name, retention time, and calculated retention indices (Lee et al. 1979). Results are given for the first two factors of the discriminant analysis describing ~ 50% of the total variance (Factor 1 and 2). (Branching points are given according to the data from Salvy et al. 2001.)

| Peak no. | Compound name | Retention time (min) | Retention index | Factor 1 | Factor 2 |
|----------|---------------|----------------------|-----------------|----------|----------|
| 2        | n-C17:0       | 14.294               | 1700            | − 0.238  | 0.060    |
| 3        | Unknown       | 14.383               | 1707            | − 0.255  | − 0.034  |
| 5        | n-C19:0       | 16.460               | 1900            | − 0.018  | − 0.283  |
| 7        | 9-C21:1       | 18.162               | 2074            | 0.078    | 0.109    |
| 8        | C21:0         | 18.397               | 2100            | 0.145    | − 0.212  |
| 9        | Unknown       | 19.015               | 2166            | 0.243    | − 0.091  |
| 11       | n-C22:0       | 19.313               | 2200            | − 0.109  | 0.004    |
| 12       | 9-C23:1       | 19.970               | 2274            | − 0.075  | − 0.333  |
| 13       | n-C23:0       | 20.192               | 2300            | − 0.081  | − 0.325  |
| 14       | 11 + 9-MeC23  | 20.484               | 2334            | − 0.258  | 0.011    |
| 15       | 7,17-diMeC23  | 20.812               | 2373            | − 0.277  | 0.056    |
| 16       | n-C24:0       | 21.013               | 2400            | 0.057    | 0.064    |
| 17       | 9-C25:1       | 21.650               | 2476            | − 0.255  | − 0.116  |
| 18       | n-C25:0       | 21.842               | 2500            | − 0.225  | − 0.081  |
| 19       | 13 + 11 + 9 + 7-MeC25 | 22.097               | 2533            | − 0.215  | 0.162    |
| 20       | 5-MeC25       | 22.228               | 2549            | − 0.145  | 0.199    |
| 21       | 4-MeC25 + 9,17-diMeC25 | 22.327               | 2562            | 0.172    | 0.180    |
| 22       | 3-MeC25 + 7,15-diMeC25 + 7,17-diMeC25 | 22.391               | 2570            | 0.155    | 0.290    |
| 23       | 5,15-diMeC25  | 22.487               | 2583            | − 0.184  | 0.078    |
| 25       | C27:2 + 14-MeC26 | 22.855               | 2631            | 0.104    | 0.231    |
| 26       | 9-C27:1 + x,y-diMeC26 | 23.186               | 2675            | − 0.041  | − 0.109  |
| 27       | n-C27:0       | 23.373               | 2700            | 0.283    | 0.004    |
| 28       | 15 + 13 + 11 + 9 + 7 + 5-MeC27 | 23.584               | 2731            | − 0.069  | 0.189    |
| 29       | 11,17-diMeC27 + 11,15-diMeC27 + 9,x-diMeC27 | 23.774               | 2756            | − 0.058  | 0.360    |
| 32       | n-C28:0       | 24.083               | 2800            | 0.259    | 0.031    |
| 33       | 14 + 13 + 12 + 11-MeC28 | 24.295               | 2829            | 0.193    | 0.182    |
| 34       | 9-C29:1 + C29:2 + 2-MeC28 | 24.634               | 2877            | 0.204    | − 0.213  |
| 35       | n-C29:0       | 24.793               | 2900            | 0.261    | − 0.047  |
| 36       | 15 + 30 + 11 + 9 + 7-MeC29 | 25.002               | 2931            | 0.018    | 0.098    |
| 40       | 9-C31:1       | 25.992               | 3080            | − 0.018  | − 0.244  |
| 41       | n-C31:0       | 26.111               | 3100            | 0.187    | − 0.125  |
| 42       | 15 + 13 + 11-MeC31 | 26.305               | 3127            | − 0.026  | 0.146    |
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samples according to their phenotypic changes. The successful infection and resulting disease-specific phenotype was characterized by a high mortality rate of infected individuals until day 21 post-grafting; corresponding well with data of previous studies (Grossar et al. 2020; Lewkowski and Erler 2019; Nakamura et al. 2016; Riessberger-Gallé et al. 2016). The second parameter for monitoring the course of the disease was larval colour change (Fig. 1). Characteristic colour change made it possible to combine larvae to different ‘groups’ of clinical symptoms accompanied with EFB disease (Forsgren et al. 2013, 2018). Finally, all larvae that survived until day 8 showed strong differences in body weight between infected and control larvae. *M. plutonius*-infected larvae are usually lighter, compared to control individuals (Bailey 1960, 1963; Lewkowski and Erler 2019). This difference in larval weight is attributed to starvation (Tarr 1937), driven by the host–pathogen competition for nutrients. Under natural conditions, honey bee larvae receive their food from nurse worker bees by progressive feeding and worker bees modify food composition depending on larval developmental stage (≤ 3d: royal jelly, > 3d: worker jelly). This means worker larvae have limited access to food jelly, they do not ‘float’ on huge quantities of food jelly as they did under artificial rearing in the current study. The higher food demand, solicited from starving infected larvae, may not be compensated by higher feeding frequency of worker bees in the hive. Consequently, the weight of infected larvae might be reduced even stronger under natural conditions as symptom of the bacteria-caused malnutrition of the larvae (Bailey 1959, 1960; Lewkowski and Erler 2019; Nakamura et al. 2016). Nevertheless, larval weight can be used as valid criterion to characterize the course of the disease under controlled in vitro rearing conditions.

While colour, weight and survival are mostly visual parameters describing the disease, worker bees use olfactory cues for orientation and brood care inside the hive. Removing diseased brood is the second step of hygienic behaviour after successful identification and differentiation from healthy individuals. Olfactory marker substances or signature mixtures are well known to initiate hygienic behaviour. For example, Swanson and colleagues (2009) detected three specific compounds (phenethyl acetate, 2-phenylethanol and benzyl alcohol) in 5-day-old larvae infected by the fungal pathogen *Ascosphaera apis* (chalkbrood), and only one single compound (phenethyl acetate) induced hygienic behaviour on chalkbrood diseased individuals or paraffin larval dummies treated with phenethyl acetate. 2-Phenylethanol was detectable in trace quantities in 20% of our EFB-samples, however without any disease-specific pattern. Day 7 old honey bee larvae, infected with *P. larvae* (AFB pathogen), were characterized by several volatile disease markers (propionic acid, valeric acid, 2-nonanone) that can be detected by more sensitive middle-aged bees rather than by forager bees.
Lee et al. (2020). However, it is still elusive if and how they are involved in hygienic behaviour. Phenethyl acetate and the volatile diagnostic substances of American foulbrood disease were not present in diseased larvae of the current study. Benzyl alcohol and propionic acid were not detectable in *M. plutonius*-infected larvae as both run in parallel with the solvent peak due to their low Kovats-indices. Solid phase micro extraction (SPME) can be used in future studies to investigate highly volatile substances (e.g., H2S) that might also be involved in the eponymous foul smell of the brood. Currently, with the random presence of 2-phenylethanol we might conclude that the fungal disease chalkbrood and the bacterial diseases (AFB and EFB) do not share the same identification pattern for hygienic worker bees. Even volatile disease markers, specific for another bacterial brood disease (AFB), that have the potential to be chemical cues for worker bees, could not be detected for EFB-disease. Altogether, we see that each microbial infection results in very distinct chemical profiles and there is no general disease signal for sick honey bee larvae. However, we have to mention that most studies focused mainly on the identification of disease-specific diagnostic substances. The identification of unique compound(s) in diseased brood is only the first step in determining biological relevance. Future studies should combine both, the characterization of the disease-associated identification pattern among the range of CHCs and BEPs, and the identification of putative signal substances. Hygienic bees may simply recognize the deviation from ‘normal’ by comparing the scent of healthy and diseased brood, which is the signal (pattern deviation) to eliminate ‘abnormal’ individuals. The innate scent of healthy brood can be expected to be learned based on the different tasks performed along the age polyethism during individual development of worker bees. In the case of EFB, worker bees might be even stricter when food resources become scanty.

Analysing CHC profiles, a relatively high proportion of the total variance, in addition to the first dimension (larval age), can be explained by the second dimension of the CHC discriminant analysis (Fig. 3); dividing the dataset into infected and uninfected individuals. *M. plutonius* infection caused a change in the composition of larval CHC profiles. The main compounds characterizing the treatment-dependent CHC change were odd hydrocarbons.

**Fig. 3** Discriminant analysis of honey bee larvae cuticular hydrocarbon profiles separated by age (5 vs. ≥ 8 days post-grafting) and treatment (infected vs. control), by the two first dimensions (49.9% of the total variance). Sample labels contain larval age (day post-grafting), treatment (infected—triangles, control—circles) and colour (w white, y yellow, y/br yellow/brown, br brown, bl black) according to Table 1 and Fig. 1. Ellipses highlight distinct sample groups.

**Fig. 4** Discriminant analysis of honey bee larvae brood pheromone profiles with a clear cluster (encircled) for older infected larvae, surrounded by older healthy and young (day 5) healthy and infected larvae. Sample labels contain larval age (day post-grafting), treatment (infected—triangles, control—circles) and colour (w white, y yellow, y/br yellow/brown, br brown, bl black) according to Fig. 1.
Changes in chemical cues of *Melissococcus plutonius* infected honey bee larvae

As cues modulating feeding and/or brood removal behaviour, the total variance was caused by age-dependent differences in 

*Varroa destructor* and *Varroa jacobsoni* infected honey bee pupae (Martin et al. 2001, 2002; Salvy et al. 2001; Wagoner et al. 2019). Changes in characteristic CHCs that are candidates for hygienic signalling were also stock-specific (Wagoner et al. 2019), therefore several hygienic lineages have to be tested to identify marker substances for hygienic selection. In the same context two substances have been identified recently, (Z)-6-pentadecene (C\textsubscript{15}) as biologically activity substance, and (Z)-10-tritriacontene (C\textsubscript{33}) as a novel hygienie trigger (Wagoner et al. 2020). *Varroa destructor* infected (pre)-pupae further showed comparable results for BEP compounds with distinct BEP profiles for removed brood and differences in amounts of BEP components, in particular, of removed pupae (Mondet et al. 2016). Hygienic worker bees might use these disease-specific cues to identify infested brood cells (Mondet et al. 2016). In this study, older *M. plutonius*-infected larvae exuded a specific BEP bouquet, a potential chemical cue for brood removal. Once again, we want to stress our alternative hypothesis that hygienic workers may not use diagnostic substances or cues, they may simply remove everything that does not smell ‘normal’. This seems plausible as worker bees do not need to ‘learn’ new substances for each disease; they only need to compare the odorant bouquet with that of normal, healthy brood. Potential mechanisms of diseased brood odorant bouquet comparison, in the olfactory lobes and the brain, against the excitation pattern of normal brood, and subsequent brood removal if patterns are non-matching, is still elusive and has to be tested in future experiments.

*Brood pheromone blends, even single compounds, are well known to provide context-specific signals that induce different behaviours* (Slessor et al. 2005). However, they need further investigation in the context of EFB disease, to understand their potential role for infected brood removal, as shown for *Varroa*-infested bees. Biotests with healthy or diseased brood/dummies, by manipulating BEP composition and quantities, might be a future project to unravel the actual signal for hygienic brood removal. However, it is mandatory to use different concentrations, as pheromone response ranges can be quite narrow and compound specific (Le Conte et al. 1995).

The CHCs discriminant analysis of this study, along with strong differences for β-ocimene, showed that 34.5% of the total variance was caused by age-dependent differences in the composition of volatile and non-volatile substances emitted by larvae. These age-related differences may act as cues modulating feeding and/or brood removal behaviour. On the contrary, age-related changes of CHCs, BEPs, and β-ocimene might be hard to differentiate from disease-related changes as the disease and its symptoms develop gradually over time. During development, the malnutrition of larvae as symptom of EFB disease might cause a higher emission of β-ocimene. Such a variation may be recognized by the nurse bees, in relation to a reference (e.g., BEP), and may trigger hygienic behaviour of workers. Here, under non-natural in vitro conditions, β-ocimene was detected in all larval stages with higher values for young individuals (day 5). It seems that the observed result is contrary to our expectation that older larvae show higher β-ocimene values than younger. However, the in vitro rearing may prevent the production of high quantities of β-ocimene. The artificial rearing protocol provides worker larvae with essential nutrients for completing their development, but their food consumption is limited for the 6-day feeding period (Crailsheim et al. 2013). This may result in the production of higher amounts of β-ocimene for the younger larvae to beg for additional food, based on the empty pharynx of the larvae. Infected larvae and larvae not completing their larval phase until day 7/8 are expected to increase starvation signalling via β-ocimene production. Contrary to this expectation, infected as well as older (non-pupa) larvae produced lower quantities of β-ocimene, which might be explained by the presence of non-consumed food that accumulated over time. The availability of food will result in absence of signaling of an empty pharynx. Consequently, begging for additional food was not necessary. However, this mechanism will not prevent the inefficient uptake of nutrients in the gut system for larval metabolism. Finally, larvae floating on non-consumed food will starve due to ineffective nutrient uptake even though they had enough food.

In natural EFB disease outbreaks, food limitation is of major importance. If food stores become empty and successful foraging does not ensure sufficient food availability in the bee hive, worker bees are supposed to eject more larvae (Bailey 1960). In this and other cases of insufficient food provisioning of larvae, β-ocimene is the relevant signal to communicate with the nurse bees. A potential reference substance or blend for the worker bees to estimate the signal strength of β-ocimene has to be discovered in future experiments. BEP, C\textsubscript{25} or C\textsubscript{27} quantities are potential candidates.

Infected older larvae (≥ day 8) were the only individuals in the current study that produced the necromone (death pheromone) oleic acid that is known as ‘ejection’ signal, which may induce infected brood removal. Various other studies showed that sick or dead larvae produce irregular high amounts of pheromones supposed to trigger hygienic behaviour. For instance, 12–17-day-old freeze killed pupae emitted significantly more β-ocimene and oleic acid (McAfee et al. 2017). Oleic acid has a central function for hygienic brood removal of *Varroa*-infested and dead pupae, and potentially for *P. larvae* infected brood (Lee et al. 2020;
Martin et al. 2002; McAfee et al. 2017, 2018). However, if worker bees use oleic acid as signal to removed EFB-infected brood has to be confirmed by bioassay of future studies.

Comparing the chemical bouquet of healthy and diseased larvae, we discovered a so far unknown compound in the honey bee A. mellifera. M. plutonius-infected larvae extracts contained γ-octalactone, and quantities increased with increasing M. plutonius loads. So far, only few studies detected γ-octalactone (alias γ-octanoic lactone) in cephalic extracts from workers of A. laboriosa (Blum et al. 2000) and in the airspace around A. dorsata bees captured in vials (Li et al. 2014). The authors concluded that γ-octalactone might possess a specific pheromone that elicit significant alarm response (Blum et al. 2000; Li et al. 2014). γ-Octalactone-specific avoidance response has also been shown for A. cerana forager bees, for a compound that is not found in A. cerana alarm pheromone (Wang et al. 2016). This shows that γ-octalactone might also be produced by diseased honey bee larvae that may signal illness to workers and may induce hygienic behaviour. For future studies, γ-octalactone can be an ideal marker signal for diseased brood, even before visible signs can be detected. Nevertheless, the potential function of γ-octalactone for EFB disease-associated hygienic behaviour remains unknown.

Other cues for detecting M. plutonius infected larvae might originate from the bacterium itself, since microorganisms may produce taxa or even strain characteristic compounds like specific metabolites (Djukic et al. 2018; Lee et al. 2020). Bacteria derived disease markers (e.g., propionic acid and valeric acid) have recently been identified for P. larvae infected honey bee larvae (Lee et al. 2020). Different strains may also have various impact on larval odorant bouquet just as they have on larval mortality (Grossar et al. 2020; Lewkowski and Erler 2019; Nakamura et al. 2016). An analysis of the food samples revealed that there is a difference in composition of hydrocarbons between food remains of infected and control larvae, maybe caused by M. plutonius (Fig. S4). However, food samples were contaminated by larval faeces, which might have increased M. plutonius concentrations in samples of the infection group, leading to very distinct GC–MS profiles. Under natural hive conditions, further approach is given by secondary invaders, which are relevant to the content of this article.

In conclusion, we demonstrated that M. plutonius infection-dependent changes of the odorant bouquet of artificially reared and infected A. mellifera larvae are detectable by GC–MS. The comparison of sick and healthy individuals revealed that γ-octalactone, brood ester pheromones, β-ocimene, oleic acid and other substances, mostly long-chained hydrocarbons, are a good indication for an EFB-specific key for worker bees. These are prime candidates for EFB-associated hygienic behaviour, which does not match with key substances of other diseases, and clearly demonstrates that there is no unique signal to remove sick brood. The compounds identified in the current study might not only be relevant for EFB disease-specific brood removal, they have also been demonstrated to strongly act on honey bee worker behaviour in general (kin and nestmate recognition, brood recognition, controlling feeding behaviour, in-hive orientation, etc.). However, a direct link between emitted EFB disease-associated chemical bouquets and hygienic behaviour is still pending. A field bioassay; i.e., application of γ-octalactone (and possibly in combination with other compounds such as BEPs, oleic acid and β-ocimene) to healthy brood in field colonies is needed to determine if the compounds elicit hygienic behaviour. Furthermore, a direct link between EFB disease and hygienic behaviour, especially in field, is still missing (only Bailey 1960) and has never been studied in natural disease outbreaks. Nevertheless, with the known function of olfactory signals for the honey bee superorganism, we hypothesize that the results of this study show to have an effect on the chemical interaction of honey bee brood, healthy or diseased, and hygienic workers bees.

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Data availability Raw data are available within the supplementary material.

Compliance with ethical standards

Conflict of interest The authors have no conflicts of interest to declare that are relevant to the content of this article.

Ethics approval Endangered or protected species were not used in this study. Experiments and observations conform to the laws of Germany in relation to animal protection. No specific ethics certification was required for this research.

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