Formic acid modulates latency and accuracy of nestmate recognition in carpenter ants

David Baracchi$^{1,2*}$, Martin Giurfa$^{1,3}$, Patrizia d'Ettorre$^{3,4}$

$^1$Research Centre on Animal Cognition, Center for Integrative Biology, CNRS, University of Toulouse, France

$^2$Department of Biology, University of Florence, Sesto Fiorentino, Italy

$^3$Institut Universitaire de France (IUF)

$^4$Laboratory of Experimental and Comparative Ethology, University Sorbonne Paris Nord, France

* To whom correspondence should be addressed. E-mail: david.baracchi@unifi.it

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**Summary statement**

Exposure to an alarm pheromone increases both latency and accuracy of the response to recognition cues in ants.
Abstract

Decision-making processes face the dilemma of being accurate or faster, a phenomenon that has been described as speed-accuracy trade-off (SAT) in numerous studies on animal behaviour. In social insects, discriminating between colony members and aliens is subjected to this trade-off as rapid and accurate rejection of enemies is of primary importance for the maintenance and ecological success of insect societies. Recognition cues distinguishing aliens from nestmates are embedded in the cuticular hydrocarbon (CHC) layer and vary among colonies. In walking carpenter ants, exposure to formic acid (FA), an alarm pheromone, improves accuracy of nestmate recognition by decreasing both alien acceptance and nestmate rejection. Here we studied the effect of FA exposure on the spontaneous aggressive mandible opening response of harnessed *Camponotus aethiops* ants presented with either nestmate or alien CHCs. FA modulated both MOR accuracy and the latency to respond to odours of conspecifics. In particular, FA decreased MOR towards nestmates but increased it towards aliens. Furthermore, FA decreased MOR latency towards aliens but not towards nestmates. As response latency can be used as a proxy of response speed, we conclude that contrary to the prediction of the SAT theory, ants did not trade off speed against accuracy in the process of nestmate recognition.
Introduction

The recognition of group members is important for the evolution of cooperation and the maintenance of social life (Hamilton 1987). In social insects, discriminating colony members from aliens allows to direct appropriately altruistic behaviours without incurring in the cost of cooperating with intruders. Moreover, recognizing and reacting promptly to social parasites, robbers or predators is vital for colony success. As a result, insects living in social groups typically excel in discriminating friends and foes (d’Ettorre and Lenoir 2010). Social recognition systems are based on a multitude of cues from different sensory modalities among which vision and olfaction play a significant role (Tibbetts 2002; van Zweden and d’Ettorre 2010; Baracchi et al. 2016). In ants, recognition systems are predominantly based on the layer of hydrocarbons coating the cuticle of individuals, which defines the chemical signature of colonies (d’Ettorre and Lenoir 2010; Bos and d’Ettorre 2012). Cuticular hydrocarbons (CHCs) constitute a blend of many chemical compounds, mainly linear alkanes, alkenes and methyl-branched alkanes (van Zweden and d’Ettorre 2010), which vary qualitatively among different species, and quantitatively among colonies of the same species, or even among individuals belonging to different morphological or physiological castes (Vander Meer and Morel 1998; Monnin 2006). The sophisticated olfactory system of ants detects CHCs at very short distance (Brandstaetter et al. 2008) and resolves up to the individual level the identity of opponent ants (D’Ettorre and Heinze 2005), securing the nest from exploiters.

CHCs are not the only chemical cues that mediate social interactions in ants and other social insects. Volatile pheromones are also used to alert colony members to coordinate their defence against exploiters (Blum 1969; Nouvian et al. 2016). Pheromones are intraspecific chemical messengers that trigger context and signal-specific, adaptive responses (Karlson and Lüscher 1959; Wyatt 2014). Their primary function is to convey a message to
one or more receivers, eliciting thereby a fast, highly predictable, and adaptive response. Yet, pheromones are not just chemical messengers. Recent work has uncovered a novel function for these substances, namely the modulation of the subjective evaluation of reinforcing stimuli (e.g. reward or punishment). Pheromones can thus modify the responsiveness to aversive or appetitive stimuli (Urlacher et al. 2010; Baracchi et al. 2017; Rossi et al. 2018; Baracchi et al. 2020). Such a modulatory effect was also detected when Camponotus ants were pre-exposed to the alarm pheromone formic acid (FA) in the context of social interactions (Rossi et al. 2019). In this case, pheromone pre-exposure improved nestmate discrimination accuracy by increasing aggressive behaviours towards aliens, while decreasing simultaneously aggression erroneously directed towards nestmates (Rossi et al. 2019).

Although the exact neural mechanisms underlying this modulatory action of FA remain to be elucidated, it was suggested that this pheromone modulates attentional processes and thus the sensitivity to recognition cues (Rossi et al. 2019).

Although accuracy is certainly a crucial aspect of any recognition process, the speed of recognition is equally important (Heitz 2014). Both variables are intimately connected in many decision-making processes (Wickelgren 1977; Heitz 2014) and may operate in orthogonal ways as decision accuracy depends on being well informed, which requires time. On the contrary, being faster in a decision process could occur at the expense of being accurate. The relationship between speed and accuracy in decision making has been refereed as the speed–accuracy trade-off (SAT) (Busemeyer and Townsend 1993). Examples of SAT have been described for several social insect species, in many different ecologically relevant tasks, including foraging, predator detection, prey choice and communication (Wickelgren 1977; Franks et al. 2003; Ings and Chittka 2008; Trimmer et al. 2008; Chittka et al. 2009).

For instance, when foraging bumblebees were tested in a colour discrimination task, some
bees made rapid choices but with low precision, while other bees were slower but highly
accurate (Chittka et al. 2003).

Here we focused on nestmate recognition in carpenter ants and studied whether the
alarm pheromone FA affects not only the accuracy (Rossi et al. 2019) but also the speed of
this process in an attempt to determine to what extent pheromones act on SAT processes. To
this end, we pre-exposed individually harnessed ants to FA and quantified afterwards their
mandible opening response (MOR) to a glass rod coated with alien or nestmate CHCs
(Guerrieri and d'Ettorre 2008). This stereotyped defensive response has been already used to
study both within- and between-species aggression in various ant species and aversive
associative learning of carpenter ants (Desmedt et al. 2017). We thus determined if the speed
and the accuracy of the recognition process are traded off and affected by pheromone pre-
exposure.

Material and Methods

Study Species and Housing

We used four queen-right colonies of *Camponotus aethiops* (Latreille 1798) collected in 2016
at Pompertuzat (Midi-Pyrénées, France). Colonies were kept under controlled laboratory
conditions (25°C, light-dark cycle = 12:12, ~ 50% relative humidity). Each colony was
housed in a plastic box (26 × 19 × 10 cm) with plaster floor connected by a tube to another
box conceived as a foraging arena (26 × 19 × 10 cm) containing sand on the floor. The nest
box was covered by cardboard in order to make it dark while the foraging arena was exposed
to light. The inner faces of the two boxes were coated with Fluon® (AGC Chemicals Europe,
Thornton Cleveleys, Lancashire, UK) to prevent ants from escaping. Ants were fed twice a
week with pieces of crickets and flour worms for proteins and honey/apple mix for
carbohydrates and vitamins. Water was provided *ad libitum*.
Nestmate recognition assay

We designed an experiment to determine whether the alarm pheromone formic acid (FA, analytical grade, Sigma-Aldrich) modulates the latency and the accuracy of the responsiveness to nestmate and non-nestmate odours. On each experimental day, medium size forager ants were gently collected with tiny forceps from the foraging arena of one of the four colonies. Ants were immediately cold anesthetized on crushed ice for a few minutes and individually harnessed in small plastic holders. A small strip of adhesive tape between the head and the thorax was used to immobilize the ants, so that they could only freely move their antennae and mouthparts (Guerrieri and d'Ettorre 2008). Once harnessed, ants were kept resting for three hours in a dark and humid place at room temperature (about 60-70% relative humidity, 24 ± 2°C) to let them acclimatize to the new restraining situation. After resting, ants were randomly allocated to either the control group and exposed to pure water (solvent) or to the experimental group and exposed to FA.

A first assay (test 1) was performed before exposure to quantify basal responsiveness to nestmate and non-nestmate CHCs. Ant responsiveness to these chemicals was quantified using the mandible opening response (MOR), (Guerrieri and d'Ettorre 2008). The test entailed eight presentation trials: four nestmate trials (A) and four non-nestmate trials (B) in a pseudorandom sequence, such as ABABBABA, so that the same stimulus (A or B) was never presented more than twice consecutively. A 12-minute inter-trial interval was used. During each trial, one ant at a time was placed under a stereomicroscope (Leica S8 APO, magnification 10 ×) in order to better visualize its MOR. Each trial lasted 25 s and consisted of 10 s of familiarization with the experimental context, 10s of stimulus (nestmate or non-nestmate CHCs) presentation and 5 s of post-stimulus resting in the setup. Each chemical stimulus was presented to the harnessed ant on a glass rod whose tip was previously coated with the CHC extract of either nestmate or non-nestmate ants (see below). The glass
rod was carefully manoeuvred by means of a micromanipulator (WPI, M33) to avoid contaminations. Upon stimulation, the rod was placed always at the same distance (2 mm from the head) of the antennae. Each stimulus was preceded by the presentation of a clean rod (presented by hands) in order to familiarize the ants with the visual component of this stimulus.

Fifteen min after the end of this first assay (test 1), ants were exposed either to FA (experimental group) or to the solvent alone (pure water, control group) to determine if pheromonal exposure modified their MOR responses. To this end, harnessed ants were individually confined for 15 min in a 50 ml plastic bottle containing a filter paper (1 x 5 cm) soaked either with the pheromone or pure water (Rossi et al. 2019). The entire procedure was performed under a hood. FA was diluted to 12% (3 μl pheromone + 22 μl water, equivalent to one third of the content of one poison gland (Stumper 1952). Control ants were exposed to 25 μl of water. After exposure, ants were kept resting for additional 30 min (Rossi et al. 2019) and then tested again (test 2) for responsiveness to nestmate and non-nestmate odours using the same procedure as in test 1.

CHC extracts were obtained by washing pools of 5 nestmate or non-nestmate ants in 2.5 ml of solvent (pentane, HPLC grade, Sigma Aldrich) for 10 min (Rossi et al. 2019). The amount of nestmate and non-nestmate odour used in each presentation was equivalent to that of a single ant. The tips of the rods were coated by adding drops of the chemical extracts using a micropipette and the rods were let dry for 1 h before starting the experiment to ensure that the solvent (pentane) evaporated. To avoid real replicates during the eight presentation trials within each assay (test 1 and test 2), alien and nestmate extracts were obtained from 4 pools of alien and nestmate ants, respectively. In the case of non-nestmates, each pool belonged to a different colony. Each presentation was video recorded from above with an integrated microscope camera. The latency to display the MOR from the moment in which
the rod was positioned at 2 mm from the head and the occurrence of MOR (yes/no) to each
stimulus presentation were quantified.

Locomotor activity assay

In order to determine whether FA merely affected motor responses, thus influencing the
observed MOR results, we designed a simple assay to monitor the locomotor activity of free
walking ants, pre-exposed either to FA or to water, which is described in the Supplementary
Materials. The results show that FA did neither impair nor modulate the locomotor activity of
ants (Fig. S1).

Data analysis and statistics

To study the effect of FA in terms of population response the proportion of reacting ants and
the speed of their response to nestmate and non-nestmate stimulations, after and before
exposure, were analysed using ANOVA designs. For testing accuracy, individual ant
responses (MOR: 1 or 0) were examined using generalized linear mixed models (GLMMs)
with a binomial error structure - logit-link function - , glmer function of R package lme4
(Bates et al. 2014). The speed of the response was analysed using GLMMs fit with Poisson
family distribution and identity link function. Q–Q plots and scatterplots of the residuals of
the model were checked visually for normal distribution and homoscedasticity. In both cases,
independent analyses were performed for ants exposed to water and ants exposed to FA. In
all the models ‘Ant response’ was entered as dependent variable, “Treatment time”
(before/ after exposure to either water or FA) and “Odour stimulus” (nestmate/non-
nestmate extract) as fixed factors, and “Trial” as covariate. Moreover, ‘Individual identity’
(IDs) was considered as a random factor to allow for repeated-measurement analysis. Colony
of origin was also entered as random factor. When necessary, models where optimized with
the iterative algorithms BOBYQA or Nelder-Mead. In each analysis, several models were run
and compared to identify significant interactions between fixed factors and/or covariates and
the significant model with the highest explanatory power (i.e. the lowest AIC value) was
retained. Interactions, wherever significant, are reported in the text. Tukey’s post-hoc tests
were used to detect differences between the different groups (lsmeans function from R
package lsmeans (Lenth and Lenth 2018).

To study the effect of FA at the individual level, for each tethered ant we calculated
a nestmate and a non-nestmate MOR Score (MS). The former was quantified as the sum of
MORs to the four nestmate presentations while the latter was quantified as the sum of MORs
to the four non-nestmate presentations. Thus, both MSs could range from 0 to 4. In the case
of nestmates, higher MS values correspond to incorrect responses (i.e. aggressive display
towards a nestmate). On the contrary, in the case of non-nestmates, higher MS values
 correspond to correct responses (i.e. aggressive display towards an alien). MSs were
calculated for test 1 (before exposure) and for test 2 (after exposure) and compared by means
of Wilcoxon signed-rank tests. We also calculated a Latency Score (LS) for each individual
ant presented with nestmate and with non-nestmate CHCs. In the case of nestmates, the LS
corresponded to the mean latency of aggressive responses upon the four nestmate odour
presentations while in the case of non-nestmates, the LS corresponded to the mean latency of
aggressive responses upon the four non-nestmate odour presentations. Both LSs were
calculated for test 1 (before exposure) and for test 2 (after exposure) and compared by means
of a Wilcoxon signed-rank test.

Finally, to test for the existence of a latency vs. accuracy trade-off in nestmate and
non-nestmate recognition, Spearman rank tests were used to correlate MSs for nestmates and
non-nestmates with nestmate and non-nestmate LSs respectively after and before FA/Water
exposure. All statistical analyses were performed with R 4.0.3 (Team 2020).
Results

In natural conditions, medium-size forager ants are typically aggressive towards alien ants and tolerant towards nestmates (Larsen et al. 2016). In the laboratory conditions in which the MOR bioassay was performed, harnessed ants reproduced this behaviour and displayed MOR to alien CHCs. In test 1 (before pheromone exposure), both FA and Water (control) groups, which were in principle identical at this point, reacted more aggressively towards the four non-nestmate presentations than to the four nestmate presentations (Fig. 1AB: within each ant category – “FA exposed” and “Water exposed” – compare the two proportions labelled as ‘Before’). This difference in the proportion of ants responding to either odour was significant (GLMM, Water group: Odour stimulus: $\chi^2 = 5.63$, df = 1, $p = 0.018$; FA group: Odour stimulus: $\chi^2 = 5.93$, df = 1, $p = 0.015$, Fig. 1AB and Fig. S2). Pheromone exposure induced a change in the proportion of ants responding with MOR to nestmate and non-nestmate odours (GLMM, Odour stimulus * Treatment time: $\chi^2 = 36.35$, df = 1, $p < 0.0001$, Fig. 1A and Fig. S2). In particular, FA exposure decreased erroneous MOR towards nestmates (GLMM, Tukey post-hoc test: $Z = -6.05$, $p < 0.0001$, Fig. 1A) while it increased correct MOR towards aliens, albeit in a non-significant way (GLMM, Tukey post-hoc test: $Z = 2.32$, $p = 0.09$, Fig. 1A). On the contrary, when ants were exposed to water, the proportion of individuals responding to nestmates and to aliens did not vary significantly (GLMM, Odour stimulus * Treatment time: $\chi^2 = 1.21$, df = 1, $p = 0.27$, Fig. 1B and Fig. S2).

In order to evaluate interindividual variability, we analysed responses in terms of individual MOR scores (MSs), which were computed both for responses to nestmate CHCs (i.e. the sum of responses to the four nestmate trials) and to non-nestmate CHCs (i.e. the sum of responses to the four non-nestmate trials). Fig. 1C shows that FA exposure significantly decreased responses to nestmates, (Wilcoxon test, $n = 69$, $V = 108$, $p < 0.0001$, Fig. 1C) and,
to a lower extent, increased responses to non-nestmates, (Wilcoxon test: n = 69, V = 961, p = 0.054, Fig. 1C). On the contrary, exposure to water did not affect the individual MS, neither towards nestmates nor to aliens (Wilcoxon test, nestmates: n = 73, V = 805, p = 0.76; alien: n = 73, V = 572, p = 0.13, Fig. 1D).

Figure 1: (A-B) Interaction plots of fitted means for the factors “Treatment time” (before/after exposure to either FA or water) and “Odour stimulus” (nestmate/non-nestmate odours). (A) MOR was differently affected by FA exposure depending on the nature of the stimulus presented so that it increased towards non-nestmate odour (GLMM, Tukey post-hoc test: p = 0.09) and it decreased towards nestmate odour (GLMM, Tukey post-hoc test: p
Ants exposed to water did not change their responsiveness neither towards nestmates nor to aliens (GLMM, *Odour stimulus* * Treatment time: p = 0.27. (C-D) Nestmate and non-nestmate MOR Score (MS) of individual tethered ants exposed either to FA or water. Boxes represent median, quartiles and max and min (upper and lower whiskers) MS values. Grey dots represent individual ants. (C) FA exposure tended to increase the MOR score (MS) to non-nestmates (p = 0.054) while it decreased it to nestmates (Wilcoxon test, p < 0.0001). (D) Water exposure did not affect the ants’ MS neither towards nestmates (p = 0.76) nor to aliens (p = 0.13).

At the population level, pheromone exposure induced a change in the mean latency of the MOR elicited by nestmate and non-nestmate odours (GLMM, *Odour stimulus* * Treatment time: χ² = 555.4, df = 1, p < 0.0001, Fig. 2AB and Fig. S3). In particular, ants exposed to FA had a shorter MOR latencies towards alien CHCs (GLMM, *Tukey post-hoc test: Z = -23.75, p < 0.0001, Fig. 2A) but did not change the MOR latency to nestmate CHCs (GLMM, *Tukey post-hoc test: Z = -0.98, p = 0.76, Fig. 2A). Overall, MOR latency towards alien CHCs decreased over the presentations and tests following FA exposure (GLMM, *Trial: χ² = 146.3, df = 1, p < 0.0001). On the contrary, ants exposed to water did neither change the latency of MOR towards nestmate nor to alien CHCs (GLMM, *Treatment time: χ² = 0.17, df = 1, p = 0.68; *Odour stimulus* *Treatment time: χ² = 1.01, df = 1, p = 0.31, Fig. 2B and Fig. S3). Overall, the MOR latency increased over the presentations (GLMM, *Trial: χ² = 86.1, df = 1, p < 0.0001). At the individual level, FA exposure significantly decreased the Latency Score (LS) to non-nestmates (Wilcoxon test: n = 56, V = 414, p = 0.002, Fig. 2C) but not to nestmates (Wilcoxon test: n = 31, V = 328, p = 0.12, Fig. 2C). After water exposure, ants slightly
decreased their LS towards aliens (Wilcoxon test, n = 58, V = 593, p = 0.043, Fig. 2D), but not to nestmates (Wilcoxon test, n = 52, V = 566, p = 0.37, Fig. 2D).

**Figure 2:** (A-B) Interaction plots of fitted means for the factors “Treatment time” (before/after exposure to either FA or water) and “Odour stimulus” (nestmate/non-nestmate odours). (A) MOR latency was differently affected by FA exposure in a stimulus dependent manner so that it strongly decreased when ants were presented with non-nestmate odour (GLMM, *Tukey post-hoc test*: p < 0.0001) but it did not vary when the same ants were presented with nestmate odour (GLMM, *Tukey post-hoc test*: p = 0.76). (B) After water exposure ants did neither change the latency of MOR towards nestmates nor to aliens.
(GLMM, Odour stimulus * Treatment time: p = 0.31). (C-D) Nestmate and the non-nestmate latency Score (LS) of individual tethered ants exposed either to FA or water. Boxes represent median, quartiles and max and min (upper and lower whiskers) LS values. Grey dots represent individual ants. (C) FA exposure decreased the LS to non-nestmates (Wilcoxon test, p = 0.002) but not to nestmates (p = 0.12). (D) After water exposure ants slightly decreased LS towards aliens (p = 0.043, but not to nestmates (p = 0.37).

We then combined our data about Latency Score (LS) and MOR Scores (MS) to analyse the existence of a speed versus accuracy trade-off. While the quantification of MS provides a measurement of response accuracy when the ants are confronted with alien or nestmate odours, the latency of their response informs about the potential speed of their response; typically shorter latencies are associated with faster responses and higher speed while longer latencies are associated with slower responses and slower speed.

We found that the response latency was not affected by the accuracy of both nestmate and non-nestmate odour recognition in ants before exposure. Precisely, individual nestmate MS did not correlate with nestmate LS (Spearman test, n = 113, ρ = 0.09, p = 0.35). Similarly, individual non-nestmate MS did not correlate with non-nestmate LS (Spearman test, n = 130, ρ = -0.17, p = 0.053). After exposure to FA or water, no significant correlation was found when we analysed the LSs and MSs of exposed ants for nestmates (FA: Spearman test, n = 35, ρ = -0.19, p = 0.28; Water: n = 63, ρ = -0.05, p = 0.68) and non-nestmates (FA: n = 64, ρ = -0.19, p = 0.14; Water: n = 62, ρ = 0.04, p = 0.77). Thus, ants did not trade-off these two aspects of the recognition process.

Discussion
The ability to discriminate between nestmates and intruders allows colony cohesion and nest defence in social insects (Hamilton 1987). Accuracy is certainly a crucial aspect of the action component of the recognition process. Yet, the speed of the recognition is equally important (Heitz 2014). Therefore, while the existence of SAT necessarily imposes boundaries, the recognition process is expected to be as accurate and fast as possible (Wickelgren 1977; Heitz 2014). Over the course of evolution, the sensory systems of social insects have been strongly refined to achieve this goal (Stroeymeyt et al. 2010; van Zweden and d’Ettorre 2010; Ozaki and Hefetz 2014). Pheromones participate in this process as they have been naturally selected to facilitate communication and response coordination at the colony level (Blum 1969).

Alarm pheromones, in particular, coordinate defensive responses of social groups and allow individuals to react promptly with stereotyped responses towards imminent dangers, such as the presence of enemies (Blum 1969; Nouvian et al. 2016).

In a previous study, we showed that FA, the alarm pheromone of several ant species, acts as cognitive modulator by enhancing nestmate discrimination in *Camponotus aethiops* ants, even when it is no longer present in the surroundings of the targeted ant (Rossi et al. 2019). In this case, FA exposure also increased aggressive behaviours of ants walking in an arena and confronted with aliens, while it decreased simultaneously erroneous aggression towards nestmates. Here we used a more controlled setup in which harnessed ants were exposed to CHCs of aliens or nestmates and confirmed our previous findings showing nestmate recognition improvement by FA in carpenter ants. In addition, we could evaluate the incidence of response latency in this process.

Our new results show that exposure to FA not only made ants more accurate in their aggressive responses but also modulated the latency of these responses. After FA exposure, those ants that still displayed erroneously MOR to nestmate odours, did it with the same latency. On the contrary, FA exposure reduced the latency of MOR towards non-nestmate
odours. Thus, FA appears to act as a facilitator that speeds aggression towards the right targets. Most likely, these changes in response latency are relevant in natural scenarios, where faster attacks to non-nestmates would increase the probability of colony success.

The theory of SAT (Wickelgren 1977; Heitz 2014) predicts that correct decisions take longer while fast decisions are more error prone. Although we did not quantify response speed (i.e. the speed of a triggered MOR), we measured the latency of MOR, which can be used as a proxy of MOR speed (i.e. a shorter latency corresponds to a faster response consummation, while a longer latency corresponds to a slower response consummation). Our results show that carpenter ants did not trade off speed (latency) against accuracy, both before and after exposure to the alarm pheromone FA. The observed increased accuracy was not affected by the speed of the responses, as FA exposure enhanced both the accuracy and the latency of the responses. Although a trade-off between speed and accuracy has been described in various contexts involving decision making, and in different modalities (Chittka et al. 2009), there are cases in which a correlation between accuracy and sampling time has not been found both in insects (Ditzen et al. 2003) and in mammals facing olfactory-discrimination problems (Uchida and Mainen 2003). Notably, in a study on nestmate recognition by hover wasps, no SAT between speed and accuracy was found (Baracchi et al. 2015), suggesting that in this particular context SAT may be uncommon.

The increased accuracy in nestmate recognition induced by FA exposure may be explained by an enhanced sensitivity to CHCs (Rossi et al. 2019). It has been proposed that FA increases the amount of information (e.g., the number of detected CHCs) available to the ants, thus decreasing the perceived phenotypic overlap between nestmate and non-nestmate recognition cues (Rossi et al. 2019). Changes in recognition speed, which determined changes in response latency, cannot be explained by changes in motor abilities as the general locomotor activity of ants was unaffected by exposure to the pheromone (Supplementary
Material). A possibility would be that FA affected attentional processes and enhanced motivation for the defensive task by acting on brain levels of neurotransmitters that have been associated with enhanced attention and aggressive responses. Attentional processes, similar to those described in vertebrates, have been characterized in insects both at the behavioral and neurobiological levels (Dyer and Chittka 2004; Giurfa 2004; Miller et al. 2011; van Swinderen 2011; Van Swinderen and Andretic 2011). In the fruit fly Drosophila melanogaster, visual attention for moving bars is mediated by a transient increase in a 20-30 Hz local field-potential recorded in a region of the brain called the medial protocerebrum (van Swinderen and Greenspan 2003). Current views relate dopamine levels in the insect brain with arousal levels (Van Swinderen and Andretic 2011). In consequence, attenuation of dopamine release in fly mutants attenuates the 20-30 Hz responsiveness to the visual object to be attended. On the contrary, pharmacological increase of dopamine rescues this responsiveness (Andretic et al. 2005). Thus, FA may upregulate dopamine levels in the brain, enhancing thereby attention in the context of nestmate discrimination. This hypothesis is sustained by findings on defensive responses in honey bees, which are triggered by the sting alarm pheromone component isoamyl acetate (IAA) (Nouvian et al. 2018). Exposure to IAA increases defensive responses and upregulates dopamine and serotonin levels in the bee brain. While serotonin has been directly related to aggressive responses in invertebrates (Kravitz 2000; Dierick and Greenspan 2007; Alekseyenko et al. 2010; Alekseyenko and Kravitz 2014; Alekseyenko et al. 2019), the dopamine component of the response may reflect the enhanced attention required to direct appropriately an attack that may have lethal consequences for the defender bee.

In conclusion, we found that FA improved nestmate recognition in C. aethiops by acting both on the accuracy (reducing erroneous responses) and on the latency of aggressive responses (reducing the latency of appropriate attacks). Our behavioural experiments do not
allow identifying the mechanism of action of FA and neural analyses are necessary to determine if and how exposure to FA upregulates levels of biogenic amines that have been associated with aggressive responses and with attentional processes. Future research aimed at quantifying biogenic amine levels upon FA exposure and at specifically blocking/activating biogenic amines receptor might help to shed light on the underlying mechanisms of FA action. Our findings add to new perspectives developed recently positing that pheromone functions exceed the traditional framework of intraspecific communication for which they have been selected. Pheromones do more than conveying specific messages to members of the same species. In insects, for instance, they can modulate in the long term responsiveness to relevant stimuli (appetitive, aversive) in contexts that differ from the one for which the pheromone is used as a messenger (Baracchi et al. 2017; Rossi et al. 2018; Hostachy et al. 2019; Rossi et al. 2019; Baracchi et al. 2020; Murmu et al. 2020; Oberhauser et al. 2020; Rossi et al. 2020). Further studies are needed to clarify these novel functions of pheromones as neuromodulators and to understand their implications for the functioning of recognition systems in general.

Author Contributions

DB and PdE conceived the study and designed the experiments. DB performed the experiments and carried out data analyses. DB, PdE and MG contributed to the writing of the manuscript.

Competing interests

No competing interests.

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References


Stumper R (1952) Quantitative data on the secretion of formic acid by ants. Comptes rendus hebdomadaires des seances de l'Academie des sciences 234:149


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