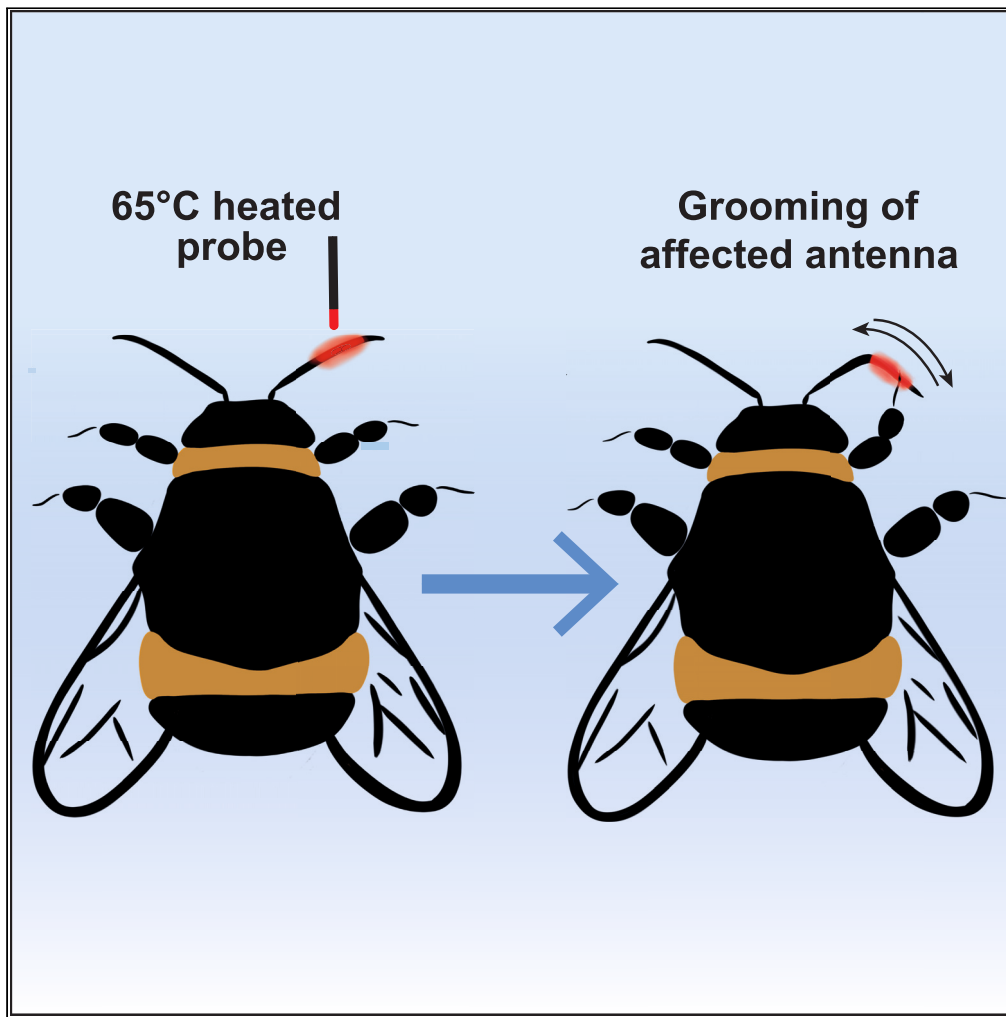


Article

Noxious stimulation induces self-protective behavior in bumblebees



Matilda Gibbons,
Elisa Pasquini,
Amelia
Kowalewska, ...,
Cwyn Solvi,
Elisabetta Versace,
Lars Chittka

matilda.gibbons@
penmedicine.upenn.edu

Highlights

We demonstrated that
bees show a form of self-
protective behavior

Bees directed grooming
toward their antenna that
was touched with a heated
probe

This self-protective
behavior occurred in the
first 2 min after stimulation

Gibbons et al., iScience 27,
110440
August 16, 2024 © 2024 The
Authors. Published by Elsevier
Inc.
[https://doi.org/10.1016/
j.isci.2024.110440](https://doi.org/10.1016/j.isci.2024.110440)

Article

Noxious stimulation induces self-protective behavior in bumblebees

Matilda Gibbons,^{1,2,8,*} Elisa Pasquini,³ Amelia Kowalewska,^{4,2} Eva Read,⁵ Sam Gibson,⁵ Andrew Crump,^{6,5} Cwyn Solvi,⁷ Elisabetta Versace,² and Lars Chittka²

SUMMARY

It has been widely stated that insects do not show self-protective behavior toward noxiously-stimulated body parts, but this claim has never been empirically tested. Here, we tested whether an insect species displays a type of self-protective behavior: self-grooming a noxiously-stimulated site. We touched bumblebees (*Bombus terrestris*) on an antenna with a noxiously heated (65°C) probe and found that, in the first 2 min after this stimulus, bees groomed their touched antenna more than their untouched antenna, and more than bees that were touched with an unheated probe or not touched at all did. Our results present evidence that bumblebees display self-protective behavior. We discuss the potential neural mechanisms of this behavior and the implications for whether insects feel pain.

INTRODUCTION

Nociception is the detection and processing of noxious stimuli¹ and can be identified from recording neural activity or behavior associated with the activation of nociceptive circuits.^{2,3} Insects have both nociceptors and nociceptive neurons that detect mechanical, thermal, and chemical noxious stimuli,^{4,5} and they respond behaviorally by moving away from and avoiding noxious stimuli.^{6,7}

Self-protective behavior—behavior with the aim to protect a body part from further noxious stimulation—is seen in response to noxious stimulation in many species, including humans. Examples include tending to, guarding, self-grooming, or rubbing a noxiously-stimulated body part. In humans, this can be seen, for example, when you grab and rub your bumped toe to reduce the pain caused by the nociceptive processing. In insects, there are no quantitative studies of self-protective behavior (such as self-grooming) directed toward a noxiously-stimulated site.⁸ In fact, anecdotal reports claim that insects do not protect their injury sites, and that insects continue to walk, feed, and mate normally after injury.^{9,10} These reports, alongside the lack of empirical evidence, are often cited as evidence against insects experiencing pain.^{11–13}

In other animals, self-protective behavior is widely reported. Rats (*Rattus norvegicus*) rub their face after being injected with a noxious substance¹⁴ and some bird species groom limbs that have been injected with a noxious substance (e.g., *Pyrrhura molinae*¹⁵). There are similar findings of fish (*Oncorhynchus mykiss*) rubbing an area that was treated with a noxious injection into the gravel and the sides of their tank.¹⁶ Some invertebrates have also been observed performing self-protective behavior, in the form of self-grooming a noxiously-stimulated site. For example, Asian shore crabs (*Hemigrapsus sanguines*) will rub a claw that has been injected with formalin.¹⁷ Similarly, shore crabs (*Carcinus maenas*),¹⁸ prawns (*Palaemon elegans*),¹⁹ cuttlefish (*Sepia pharosaensis*),²⁰ and octopuses (*Octopus bocki*)²¹ will groom or scratch a body part where acetic acid has been applied. The latter will also respond with self-grooming an area on their arm after it was squeezed with serrated forceps for up to 20 s.²²

As noted previously, there is no evidence of insects self-grooming noxiously-stimulated body parts. However, insects are known to self-groom in non-noxious contexts, for example during general cleaning,²³ and when removing dust particles (e.g., in the German cockroach *Blattella germanica*²⁴), pollen grains (e.g., in bees²⁵), and parasites such as mites (e.g., in honeybees, *Apis mellifera*²⁶). Further, after noxious stimulation, insects may also generally groom themselves more all over, or change their grooming pattern. For example, after having their antenna amputated, red mason bees (*Osmia bicornis*) groom their head and body, although, in the study where this was found, no site-specific measurements were found/taken, nor was there a non-noxious control to compare to.²⁷

Some anecdotal reports claim that insects do not protect their injury sites,^{9,10} but there are also some reports suggesting the contrary. Such reports, however, have not yet been supported by quantitative or statistical analyses.^{8,28} For example, when pinched on the abdominal

¹Department of Neuroscience, University of Pennsylvania, Philadelphia, PA 19104, USA

²School of Biological and Behavioral Sciences, Queen Mary University of London, London E1 4NS, UK

³Center for Mind/Brain Sciences, University of Trento, Rovereto 38068, Italy

⁴Academic Training Team, The Francis Crick Institute, London NW1 1AT, UK

⁵Department of Philosophy, Logic and Scientific Method, London School of Economics, London WC2A 2AE, UK

⁶Department of Pathobiology & Population Sciences, Royal Veterinary College, London NW1 0TU, UK

⁷Guangdong-Hong Kong-Macao Greater Bay Area Center for Brain Science and Brain-Inspired Intelligence, Southern Medical University, Guangzhou City, Guangdong Province 510515, China

⁸Lead contact

*Correspondence: matilda.gibbons@pennmedicine.upenn.edu

<https://doi.org/10.1016/j.isci.2024.110440>



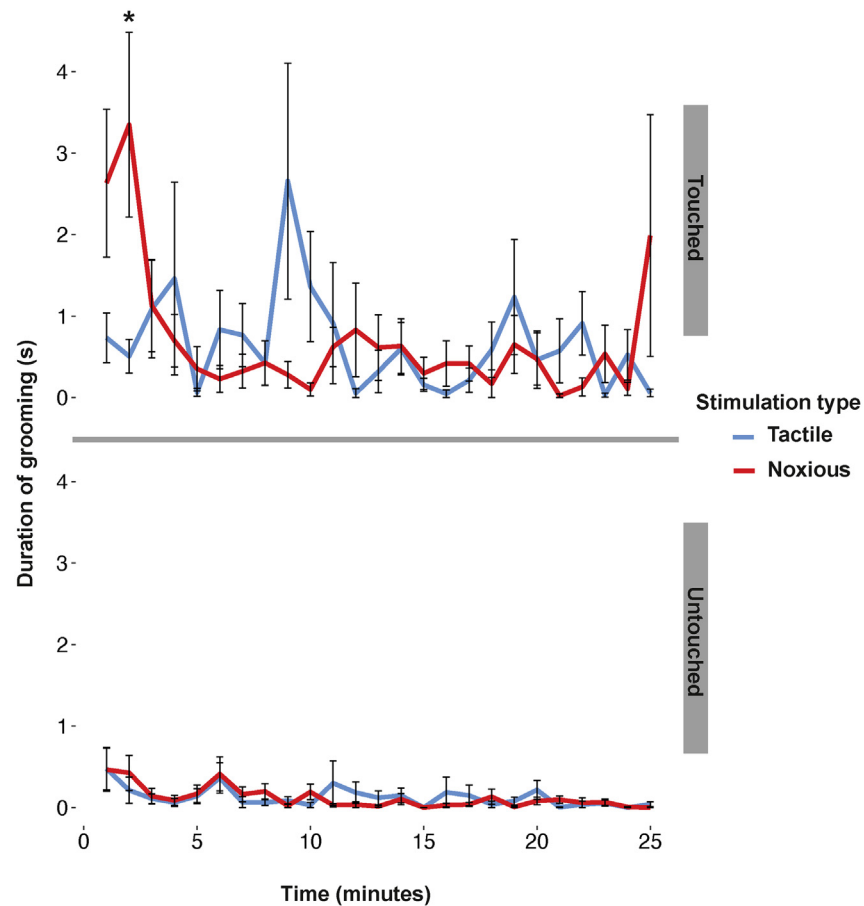


Figure 1. Mean duration of grooming for the untouched and touched antenna per each minute after noxious or tactile stimulation

* $p < 0.001$; Wilcoxon test. Error bars represent the standard error of the mean.

proleg, moth larvae (*Manduca sexta*) reportedly turn their heads to the wound, and repeatedly touch the area with their mouthparts, but this behavior was not measured or compared to a control.²⁹ The same was the case in a study on cockroaches (*Periplaneta americana*) which were reported to groom their wounds following an abdominal puncture.³⁰ Since both reports of the absence and the existence of self-protective behavior in insects are not supported by quantitative measurements or analyses,²⁸ a robust, experimental assessment of self-protective behavior in response to noxious stimuli in insects is required. The absence of quantitative empirical studies of whether insects perform self-protective behavior has fueled arguments against insects feeling pain, based on the claim that they do not protect their injury sites.^{9–13}

In this study, we tested whether *Bombus terrestris* bumblebees display a type of self-protective behavior: selectively grooming a noxiously-stimulated body part. For each bumblebee, we either briefly touched one antenna with a noxious stimulus (a 65°C heat probe), or a non-noxious tactile stimulus (an unheated probe), or we did not touch either antenna (control). We recorded self-grooming behavior toward both antennae for 25 min. If bees specifically groom a site of noxious stimulation, we would predict more grooming on the noxiously-stimulated antenna than the other antenna. We would not expect this difference in bees touched with an unheated probe, nor by bees that were not touched.

RESULTS

Self-grooming in the 25-min observation period

We first tested whether, within the whole 25-min observation period, there was a difference between grooming durations on the touched and untouched antennae, and, if so, whether this difference was larger when the probe was noxiously heated. For the 25-min observation period, bees groomed their touched antennae significantly more (touched: 18.11 ± 26.79 s; untouched: 2.22 ± 3.57 s; $t_{5792} = 5.922$; $p < 0.001$; $N = 40$), regardless of whether the stimulation was noxious or non-noxious tactile (no significant effect: $t_{5792} = 0.056$, $p = 0.955$; $N = 40$; no significant interaction: $t_{5792} = -0.224$, $p = 0.822$; $N = 40$; Figure 1). Therefore, over the 25 min, grooming was directed toward the touched antenna, but not the noxiously-stimulated antenna specifically.

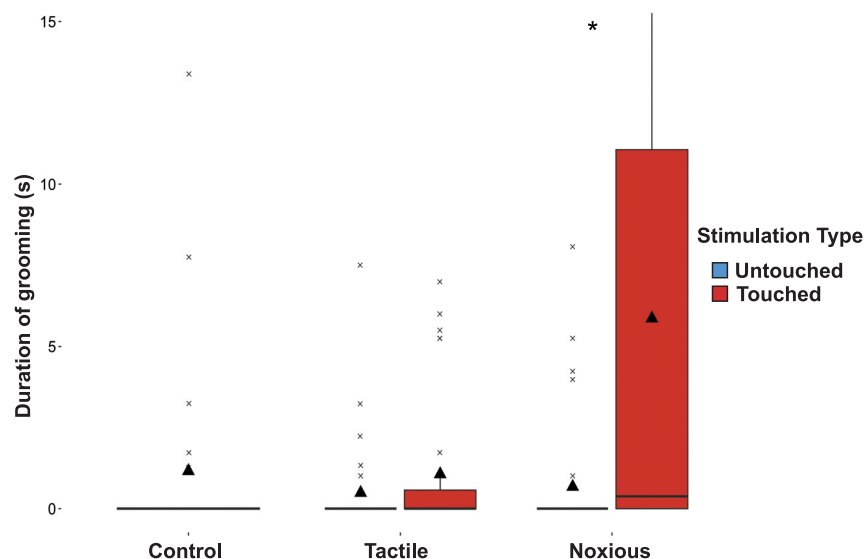


Figure 2. Boxplot of duration of grooming on each antenna for each stimulation type group

Boxplot boundaries indicate the 25th and 75th percentiles; the whiskers indicate the minimum and maximum values within 1.5 times the interquartile range. Crosses indicate values outside this range (boxplot outliers); triangles indicate the mean; lines indicate the median. * $p < 0.001$; Wilcoxon test.

We also observed a significant interaction effect of sex on the total grooming duration over the 25 min, with workers (females) grooming their touched antenna (and not their untouched antenna) for significantly longer than males (females: $N = 40$; touched antenna: 22.89 ± 30.29 s; untouched antenna: 2.60 ± 3.78 s; males: $N = 18$; touched antenna: 7.59 ± 11.33 s; untouched antenna: 1.36 ± 2.96 s; $t_{5792} = -2.665$; $p < 0.01$).

Self-grooming in the 2 mins after manipulation

In the 0–2 min time bin (the only time bin with a significant p value after applying the Holm-Bonferroni correction, see [method details](#)), bees groomed the touched antenna more than the untouched antenna when the touch was noxious (significant interaction: $t_{459} = 3.069$, $p < 0.005$; $N = 40$). This result was further supported by Wilcoxon tests: in this time bin, noxiously-stimulated bees groomed their touched antenna (6.65 ± 8.8 s) significantly more than their untouched antenna (0.75 ± 1.95 s; $W = 249.5$, $p < 0.001$; $N = 30$; [Figure 2](#)). By contrast, for tactilely-stimulated bees, there was no significant difference in grooming between the touched antenna (1.19 ± 2.23) and the untouched antenna (0.55 ± 1.57 s; $W = 324$, $p = 0.159$; $N = 28$; [Figure 2](#)). Similarly, noxiously-stimulated bees groomed for significantly longer than the tactilely-stimulated bees on the touched antenna ($t_{83} = 2.885$, $p < 0.005$; $N = 40$; [Figure 2](#)), but not on the untouched antenna ($t_{83} = 0.647$, $p = 0.519$; $N = 40$; [Figure 2](#)). There was no significant effect of sex on either the grooming in touched or untouched conditions (touched: $t_{83} = 0.253$, $p = 0.800$; $N = 40$; untouched: $t_{83} = -1.273$; $p = 0.207$; $N = 40$) nor on the overall antennal grooming in noxious and tactilely-stimulated bees ($t_{459} = -0.851$, $p = 0.395$; $N = 40$).

We then tested whether the duration of antennal grooming was greater for either the noxiously-stimulated or the tactilely-stimulated bees compared to the control (untouched) bees. Noxiously-stimulated bees groomed their touched, and not their untouched, antenna for longer than the control bees groomed either antenna (touched: 2.85 ± 5.48 s; $t_{75} = 2.55$, $p = 0.0127$; $N = 54$; untouched: 0.50 ± 1.64 ; $t_{75} = -0.318$, $p = 0.752$; $N = 54$; either antenna (mean grooming duration): 0.57 ± 2.14 ; [Figure 2](#)). There was no significant effect of sex on either the grooming compared to control in touched or untouched conditions (touched: $t_{75} = -0.111$, $p = 9.117$; $N = 54$; untouched: $t_{75} = -1.493$; $p = 0.140$; $N = 54$). By contrast, tactilely-stimulated bees did not groom either their touched or untouched antennae for significantly longer than the control bees groomed either antenna (touched: 0.77 ± 1.84 ; $t_{73} = -0.404$, $p = 0.689$; $N = 52$; untouched: 0.48 ± 1.54 ; $t_{73} = 0.228$, $p = 0.821$; $N = 52$; [Figure 2](#); either antenna (mean grooming duration): 0.57 ± 2.14). There was no significant effect of sex on either the grooming compared to control in touched or untouched conditions (touched: $t_{75} = -0.127$, $p = 0.210$; $N = 52$; untouched: $t_{75} = -1.875$; $p = 0.065$; $N = 52$).

DISCUSSION

Our results provide the first quantitative evidence of self-protective behavior in insects. In the first 2 min after noxious stimulation on an antenna, bees groomed this noxiously-touched antenna more than their untouched antenna and more than control (untouched) bees groomed either of their untouched antennae. The same results were not found in bees that were touched with a non-noxious, tactile stimulus. Further, noxiously-stimulated bees groomed their noxiously-touched antenna for longer than the tactilely-stimulated bees groomed their tactilely-touched antenna.

A significant increase in self-grooming the noxiously-stimulated antenna only being evident in the first 2 min after stimulation is consistent with studies on other invertebrates, which describe self-grooming in the first few minutes after noxious stimulation.^{17–19} A reason for this timing might be that the nociceptive processing ceased after around 2 min; this would likely change with a higher intensity of the noxious stimulus than we used here. An association between grooming and the cessation or onset of nociceptive processing has been previously noted in mice, in response to nociceptive formalin injection. There is an acute grooming phase, which apparently relates to the injection itself and lasts 3 min, then no grooming is seen for another 3 min, followed by a tonic phase that is longer-lasting and appears to correspond to formalin's inflammatory effects.^{31–33} By analogy, we suggest that, in our study, the first 2 min corresponded to an acute phase of grooming in response to the noxious heat stimulation. Based on this evidence, future research should investigate the neural processing of noxious heat stimulation in insects, and how the temporal characteristics of the self-grooming might relate.

If grooming directed toward a noxiously-stimulated antenna happens in the first 2 min after stimulation, one might expect to also find a significant increase in grooming within the first minute. Here, we did observe an increase in grooming on the noxiously treated antenna in the first minute, but this increase was not statistically significant after correcting for multiple comparisons (Figure 1). This could reasonably be explained by our use of the Holm-Bonferroni correction, which has a high risk of false negatives.³⁴

In the first 25 min after stimulation, the bees' sex had a significant effect on how long they groomed their touched antenna, regardless of whether the stimulus was noxious or not, with workers (females) grooming their touched antenna for longer on average than males did. There are currently no studies investigating sex differences in self-grooming behavior in bees, but male bees do not groom pollen off their bodies, suggesting that they might be less equipped for self-grooming in response to something touching their body.³⁵

An interesting future line of research would involve investigating the neural underpinnings of our findings. The neural processing of the noxious heat might be similar to that seen in honeybees, where nociceptive signals in the antennae are detected by thermo-sensory neurons and carried to the antennal lobe.³⁶ As for the neural circuits of self-grooming, these have, of course, only been studied in the context of general, non-noxious self-grooming. For example, research on *Drosophila melanogaster* has identified neurons in the antenna that project to the ventral brain and antennal descending neurons that, if stimulated, cause antennal grooming.³⁷ Nociceptive self-grooming in bees might use similar neural mechanisms, but more research is needed.

What might our results mean for the topic of insect pain? Firstly, we need to clarify whether and how self-protective behavior might relate to pain. Self-protective behavior has been taken as evidence consistent with the presence of pain in other animals (e.g., crustaceans,¹⁷ molluscs,^{20–22} rodents,¹⁴ birds,¹⁵ and fish¹⁶), including humans³⁸ and is included in frameworks for assessing pain in animals.^{39–41} One reason for this association is that self-protective behavior seems to reduce the feeling of pain in humans^{42,43} and is not merely a reflexive behavior. For example, self-touch has been found to reduce the painful perception of heat, even when this "heat pain" is caused by an illusion that leads participants to perceive pain without there being any nociceptive stimulus.⁴⁴ This shows that self-touch reduces pain specifically, rather than nociceptive processing.

There are, however, some historic studies in frogs and dogs with severed spines where noxious stimulation of extremities induces leg movements that are roughly directed toward the site of stimulation, suggesting that nociceptive reflexes might underlie some sort of self-protective behavior.^{45,46} However, the animal pain frameworks clarify that self-protective behavior should be directed toward the injury site^{40,47} and, in these studies, the leg movements are not directed specifically to the site of noxious stimulation. This might mean that general self-grooming in response to injury might be able to occur via nociceptive reflex loops in the spinal cord, but directing the response specifically to the site of injury may require some sort of brain processing. It should also be noted that these studies lack solid experimental measures, such as quantified behavior, mention of sample size, formal analysis, or a control experiment using non-noxious stimuli or healthy animals, so the results cannot be directly compared to our study. Moreover, it is clear that the behavior we observed requires the brain, since noxious stimulation of the antenna feeds directly into the antennal lobe of the bee brain.³⁶

In conclusion, the self-protective behavior displayed by the bees in our study both requires the brain and is akin to a behavior that is associated with pain in humans and other animals. What does this mean for the likelihood that bees can feel pain? Our study shares with others (including those on vertebrates) the challenge that it is currently impossible to be certain about whether a behavior includes the affective component of pain. Therefore, to assess whether an animal can feel pain, it is valuable to collect evidence from multiple different lines of neural, behavioral, and psychological investigations to shift probabilities for or against.^{39,40,48} Self-protective behavior is included as one of eight criteria for the evidence of pain in other animals.⁴⁰ Before our study, Adult Hymenoptera already fulfilled four of these eight indicators of pain, namely they have nociceptors⁴⁹ and sensory integrative brain regions,⁵⁰ display motivational trade-offs,⁵¹ and learn from aversive experiences.⁵² Our study thus provides evidence for a 5th criterion, self-protective behaviour, so Hymenoptera might now be considered in this framework to show "strong evidence for pain."

Therefore, our study is one of a set of studies that have found evidence of traits indicative of pain in Hymenoptera. These studies, when taken together, give reason to increase our confidence that bees may feel pain. Further, at the very least, our results are incompatible with an often-quoted argument against the existence of pain in insects—the (empirically unsubstantiated) claim that they lack any form of self-protective behavior in response to noxious stimulation.^{9–13}

Limitations of the study

We suspect that the self-grooming we observed with this set-up may only be a fraction of the bees' natural response, when not under stress or in a novel environment, since stress and novel contexts have been found to reduce the expression of behaviors after noxious stimulation in insects (honeybees⁵³), similarly to other taxa (humans⁵⁴; rodents,⁵⁵ fish,⁵⁶ birds,⁵⁷ and snails⁵⁸). The experiment contained multiple novel

and/or potentially stressful experiences and environments for the bees. For example, the stimulation itself involved them climbing onto metal forceps, being lifted out of the nest box, and immobilized during the stimulation—all potential stressors. Further, bees were isolated from the nest and other colony members during testing, and their normal route back to the nest was blocked. In future experiments, observing bees in the nest post-stimulation may lead to the identification of their more naturalistic behavior in response to noxious stimulation.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
 - Lead contact
 - Materials availability
 - Data and code availability
- EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS
- METHOD DETAILS
 - Treatments
 - Behavioral analysis
- QUANTIFICATION AND STATISTICAL ANALYSIS

ACKNOWLEDGMENTS

We thank Jonathan Birch for his insightful comments on this manuscript. This research is part of a project that has received funding from Queen Mary University of London; an Erasmus+ traineeship; Open Philanthropy grant 'Addressing key evidence gaps in the science of insect sentience'; and the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation program, grant/award number 851145.

AUTHOR CONTRIBUTIONS

M.G.: conceptualization, formal analysis, investigation, methodology, writing – original draft and writing – review and editing. E.P.: conceptualization, investigation, methodology. A.K.: investigation. E.R.: investigation and writing – review and editing. S.G.: investigation. A.C.: formal analysis, methodology and writing – review and editing. C.S.: conceptualization and writing – review and editing. E.V.: conceptualization, formal analysis, funding acquisition, methodology, supervision, and writing – review and editing. L.C.: conceptualization, funding acquisition, methodology, resources, supervision, and writing – review and editing.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: January 16, 2024

Revised: May 28, 2024

Accepted: July 1, 2024

Published: July 2, 2024

REFERENCES

1. Tracey, W.D. (2017). Nociception. *Curr. Biol.* 27, R129–R133. <https://doi.org/10.1016/j.cub.2017.01.037>.
2. Kucyi, A., and Davis, K.D. (2017). The Neural Code for Pain: From Single-Cell Electrophysiology to the Dynamic Pain Connectome. *Neuroscientist* 23, 397–414. <https://doi.org/10.1177/1073858416667716>.
3. Skljarevski, V., and Ramadan, N.M. (2002). The nociceptive flexion reflex in humans – review article. *Pain* 96, 3–8. [https://doi.org/10.1016/S0304-3959\(02\)00018-0](https://doi.org/10.1016/S0304-3959(02)00018-0).
4. Tracey, W.D., Wilson, R.I., Laurent, G., and Benzer, S. (2003). painless, a *Drosophila* Gene Essential for Nociception. *Cell* 113, 261–273. [https://doi.org/10.1016/S0092-8674\(03\)00272-1](https://doi.org/10.1016/S0092-8674(03)00272-1).
5. Maliszewska, J., Jankowska, M., Kletkiewicz, H., Stankiewicz, M., and Rogalska, J. (2018). Effect of Capsaicin and Other Thermo-TRP Agonists on Thermoregulatory Processes in the American Cockroach. *Molecules* 23, 3360. <https://doi.org/10.3390/molecules23123360>.
6. Marchal, P., Villar, M.E., Geng, H., Arrufat, P., Combe, M., Viola, H., Massou, I., and Giurfa, M. (2019). Inhibitory learning of phototaxis by honeybees in a passive-avoidance task. *Learn. Mem.* 26, 1–12. <https://doi.org/10.1101/lm.050120.119>.
7. Maliszewska, J., Marcinkowska, S., Nowakowska, A., Kletkiewicz, H., and Rogalska, J. (2018). Altered heat nociception in cockroach *Periplaneta americana* L. exposed to capsaicin. *PLoS One* 13, e0194109. <https://doi.org/10.1371/journal.pone.0194109>.
8. Veit, W. (2022). Towards a Comparative Study of Animal Consciousness. *Biol. Theory* 17, 292–303. <https://doi.org/10.1007/s13752-022-00409-x>.
9. Eisemann, C., Jorgensen, W., Merritt, D., Rice, M., Cribb, B., Webb, P., and Zalucki, M. (1984). Do insects feel pain? — A biological view. *Experientia* 40, 164–167. <https://doi.org/10.1007/BF01963580>.
10. Do Insects feel pain? (1980 (ArchiveSearch). https://archivessearch.lib.cam.ac.uk/repositories/9/archival_objects/496329.
11. Adamo, S.A. (2019). Is it pain if it does not hurt? On the unlikelihood of insect pain. *Can. Entomol.* 151, 685–695. <https://doi.org/10.4039/tce.2019.49>.
12. Stelling, T. (2014). Rethinking invertebrate pain. *N. Sci.* 221, 38–41. [https://doi.org/10.1016/S0262-4079\(14\)60381-X](https://doi.org/10.1016/S0262-4079(14)60381-X).
13. Mikhalevich, I., and Powell, R. (2020). Minds without spines: Evolutionarily inclusive animal

- ethics. *Animal sentience* 5, 1. <https://doi.org/10.51291/2377-7478.1527>.
14. Clavelou, P., Pajot, J., Dallel, R., and Raboisson, P. (1989). Application of the formalin test to the study of orofacial pain in the rat. *Neurosci. Lett.* 103, 349–353. [https://doi.org/10.1016/0304-3940\(89\)90125-0](https://doi.org/10.1016/0304-3940(89)90125-0).
 15. Paul-Murphy, J.R., Krugner-Higby, L.A., Tourdot, R.L., Sladky, K.K., Klauer, J.M., Keuler, N.S., Brown, C.S., and Heath, T.D. (2009). Evaluation of liposome-encapsulated butorphanol tartrate for alleviation of experimentally induced arthritic pain in green-cheeked conures (*Pyrrhura molinae*). *Am. J. Vet. Res.* 70, 1211–1219. <https://doi.org/10.2460/ajvr.70.10.1211>.
 16. Sneddon, L.U. (2003). The evidence for pain in fish: the use of morphine as an analgesic. *Appl. Anim. Behav. Sci.* 83, 153–162. [https://doi.org/10.1016/S0168-1591\(03\)00113-8](https://doi.org/10.1016/S0168-1591(03)00113-8).
 17. Dyuizen, I.V., Kotsyuba, E.P., and Lamash, N.E. (2012). Changes in the nitric oxide system in the shore crab *Hemigrapsus sanguineus* (Crustacea, decapoda) CNS induced by a nociceptive stimulus. *J. Exp. Biol.* 215, 2668–2676. <https://doi.org/10.1242/jeb.066845>.
 18. Elwood, R.W., Dalton, N., and Riddell, G. (2017). Aversive responses by shore crabs to acetic acid but not to capsaicin. *Behav. Process.* 140, 1–5. <https://doi.org/10.1016/j.beproc.2017.03.022>.
 19. Barr, S., Laming, P.R., Dick, J.T.A., and Elwood, R.W. (2008). Nociception or pain in a decapod crustacean? *Anim. Behav.* 75, 745–751. <https://doi.org/10.1016/j.anbehav.2007.07.004>.
 20. Kuo, T.-H., Sneddon, L.U., Spencer, J.W., and Chiao, C.-C. (2022). Impact of Lidocaine on Pain-Related Grooming in Cuttlefish. *Biology* 11, 1560. <https://doi.org/10.3390/biology11111560>.
 21. Crook, R.J. (2021). Behavioral and neurophysiological evidence suggests affective pain experience in octopus. *iScience* 24, 102229. <https://doi.org/10.1016/j.isci.2021.102229>.
 22. Alupay, J.S., Hadjisolomou, S.P., and Crook, R.J. (2014). Arm injury produces long-term behavioral and neural hypersensitivity in octopus. *Neurosci. Lett.* 558, 137–142. <https://doi.org/10.1016/j.neulet.2013.11.002>.
 23. Ringo, J.M. (2020). How do flies keep clean? Head grooming in *Drosophila*. *J. Ethol.* 38, 167–172. <https://doi.org/10.1007/s10164-019-00635-0>.
 24. El-Awami, I.O., and Dent, D.R. (1995). The interaction of surface and dust particle size on the pick-up and grooming behaviour of the German cockroach *Blattella germanica*. *Entomol. Exp. Appl.* 77, 81–87. <https://doi.org/10.1111/j.1570-7458.1995.tb01988.x>.
 25. Jander, R. (1976). Grooming and pollen manipulation in bees (Apoidea): the nature and evolution of movements involving the foreleg. *Physiol. Entomol.* 1, 179–194. <https://doi.org/10.1111/j.1365-3032.1976.tb00960.x>.
 26. Russo, R.M., Liendo, M.C., Landi, L., Pietronave, H., Merke, J., Fain, H., Muntaabski, I., Palacio, M.A., Rodríguez, G.A., Lanzavecchia, S.B., and Scannapieco, A.C. (2020). Grooming Behavior in Naturally Varroa-Resistant *Apis mellifera* Colonies From North-Central Argentina. *Front. Ecol. Evol.* 8, 590281.
 27. Rogers, L.J., Frasnelli, E., and Versace, E. (2016). Lateralized antennal control of aggression and sex differences in red mason bees, *Osmia bicornis*. *Sci. Rep.* 6, 29411. <https://doi.org/10.1038/srep29411>.
 28. Gibbons, M., Crump, A., Barrett, M., Sarlak, S., Birch, J., and Chittka, L. (2022). Can insects feel pain? A review of the neural and behavioural evidence. In *Advances in Insect Physiology* (Academic Press), pp. 155–229. <https://doi.org/10.1016/bs.aip.2022.10.001>.
 29. Walters, E., Illich, P., Weeks, J., and Lewin, M. (2001). Defensive responses of larval *Manduca sexta* and their sensitization by noxious stimuli in the laboratory and field. *J. Exp. Biol.* 204, 457–469. <https://doi.org/10.1242/jeb.204.3.457>.
 30. Hentschel, E., and Penzlin, H. (1982). Beeinflussung des Putzverhaltens bei *Periplaneta americana* (L.) durch Wundsetzung, Naloxon-, Morphin-, und Met-Enkephalingaben. *Zoologische Jahrbucher. Abteilung für allgemeine Zoologie und Physiologie der Tiere*.
 31. Mittal, N., Joshi, R., Hota, D., and Chakrabarti, A. (2009). Evaluation of antihyperalgesic effect of curcumin on formalin-induced orofacial pain in rat. *Phytother. Res.* 23, 507–512.
 32. Hunskaar, S., and Hole, K. (1987). The formalin test in mice: dissociation between inflammatory and non-inflammatory pain. *Pain* 30, 103–114. [https://doi.org/10.1016/0304-3959\(87\)90088-1](https://doi.org/10.1016/0304-3959(87)90088-1).
 33. Hunskaar, S., Berge, O.-G., and Hole, K. (1986). Dissociation between antinociceptive and anti-inflammatory effects of acetylsalicylic acid and indomethacin in the formalin test. *Pain* 25, 125–132. [https://doi.org/10.1016/0304-3959\(86\)90014-X](https://doi.org/10.1016/0304-3959(86)90014-X).
 34. Guillermou, H., Abraham, C., Annesi-Maessano, I., and Mollinari, N. (2023). Methods of Global Alpha Risk Control for Multiple Hypothesis Testing: A Pedagogical Point of View. Preprint at SSRN.com. <https://doi.org/10.2139/ssrn.4563902>.
 35. Ogilvie, J.E., and Thomson, J.D. (2015). Male bumble bees are important pollinators of a late-blooming plant. *Arthropod-Plant Interactions* 9, 205–213. <https://doi.org/10.1007/s11829-015-9368-x>.
 36. Nishino, H., Nishikawa, M., Mizunami, M., and Yokohari, F. (2009). Functional and topographic segregation of glomeruli revealed by local staining of antennal sensory neurons in the honeybee *Apis mellifera*. *J. Comp. Neurol.* 515, 161–180. <https://doi.org/10.1002/cne.22064>.
 37. Hampel, S., Franconville, R., Simpson, J.H., and Seeds, A.M. (2015). A neural command circuit for grooming movement control. *Elife* 4, e08758. <https://doi.org/10.7554/eLife.08758>.
 38. Feldt, K.S. (2000). The checklist of nonverbal pain indicators (CNPI). *Pain Manag. Nurs.* 1, 13–21. <https://doi.org/10.1053/jpmn.2000.5831>.
 39. Sneddon, L.U., Elwood, R.W., Adamo, S.A., and Leach, M.C. (2014). Defining and assessing animal pain. *Anim. Behav.* 97, 201–212. <https://doi.org/10.1016/j.anbehav.2014.09.007>.
 40. Birch, J., Burn, C., Schnell, A., Browning, H., and Crump, A. (2021). Review of the Evidence of Sentience in Cephalopod Molluscs and Decapod Crustaceans. *General - Animal Feeling*.
 41. Bateson, P. (1991). Assessment of pain in animals. *Anim. Behav.* 42, 827–839. [https://doi.org/10.1016/S0003-3472\(05\)80127-7](https://doi.org/10.1016/S0003-3472(05)80127-7).
 42. Apley, J. (1976). Pain in childhood. *J. Psychosom. Res.* 20, 383–389. [https://doi.org/10.1016/0022-3999\(76\)90090-8](https://doi.org/10.1016/0022-3999(76)90090-8).
 43. Chen, J., Abbod, M., and Shieh, J.-S. (2021). Pain and Stress Detection Using Wearable Sensors and Devices—A Review. *Sensors* 21, 1030. <https://doi.org/10.3390/s21041030>.
 44. Kammers, M.P.M., de Vignemont, F., and Haggard, P. (2010). Cooling the Thermal Grill Illusion through Self-Touch. *Curr. Biol.* 20, 1819–1822. <https://doi.org/10.1016/j.cub.2010.08.038>.
 45. E. by C. Marshall, B. Lightman, and R. England, eds. (2015). *The Papers of the Metaphysical Society, A Critical Edition* (Oxford University Press). 1869–1880.
 46. Sherrington, C.S. (1906). Observations on the scratch-reflex in the spinal dog. *J. Physiol.* 34, 1–50.
 47. Crump, A., Browning, H., Schnell, A., Burn, C., and Birch, J. (2022). Sentience in decapod crustaceans: A general framework and review of the evidence. *Animal Sentience* 7, 1. <https://doi.org/10.51291/2377-7478.1691>.
 48. Birch, J. (2017). Animal sentience and the precautionary principle. *Animal Sentience* 2, 10.
 49. Peng, G., Shi, X., and Kadowaki, T. (2015). Evolution of TRP channels inferred by their classification in diverse animal species. *Mol. Phylogenet. Evol.* 84, 145–157. <https://doi.org/10.1016/j.ympev.2014.06.016>.
 50. Strube-Bloss, M.F., and Rössler, W. (2018). Multimodal integration and stimulus categorization in putative mushroom body output neurons of the honeybee. *R. Soc. Open Sci.* 5, 171785. <https://doi.org/10.1098/rsos.171785>.
 51. Gibbons, M., Versace, E., Crump, A., Baran, B., and Chittka, L. (2022). Motivational trade-offs and modulation of nociception in bumblebees. *Proc. Natl Acad. Sci. USA* 119, e2205821119. <https://doi.org/10.1073/pnas.2205821119>.
 52. Vergoz, V., Roussel, E., Sandoz, J.-C., and Giurfa, M. (2007). Aversive Learning in Honeybees Revealed by the Olfactory Conditioning of the Sting Extension Reflex. *PLoS One* 2, e288. <https://doi.org/10.1371/journal.pone.0000288>.
 53. Núñez, J., Almeida, L., Balderrama, N., and Giurfa, M. (1997). Alarm Pheromone Induces Stress Analgesia via an Opioid System in the Honeybee. *Physiol. Behav.* 63, 75–80. [https://doi.org/10.1016/S0031-9384\(97\)00391-0](https://doi.org/10.1016/S0031-9384(97)00391-0).
 54. Ford, G.K., and Finn, D.P. (2008). Clinical correlates of stress-induced analgesia: Evidence from pharmacological studies. *Pain* 140, 3–7. <https://doi.org/10.1016/j.pain.2008.09.023>.
 55. Butler, R.K., and Finn, D.P. (2009). Stress-induced analgesia. *Prog. Neurobiol.* 88, 184–202. <https://doi.org/10.1016/j.pneurobio.2009.04.003>.
 56. Thomson, J.S., Deakin, A.G., Cossins, A.R., Spencer, J.W., Young, I.S., and Sneddon, L.U. (2020). Acute and chronic stress prevents responses to pain in zebrafish: evidence for stress-induced analgesia. *J. Exp. Biol.* 223, jeb224527. <https://doi.org/10.1242/jeb.224527>.
 57. Gentle, M.J., and Corr, S.A. (1995). Endogenous analgesia in the chicken. *Neurosci. Lett.* 201, 211–214. [https://doi.org/10.1016/0304-3940\(95\)12181-1](https://doi.org/10.1016/0304-3940(95)12181-1).
 58. Kavaliers, M. (1987). Evidence for opioid and non-opioid forms of stress-induced analgesia in the snail, *Cepaea nemoralis*. *Brain Res.* 410,

- 111–115. [https://doi.org/10.1016/S0006-8993\(87\)80029-X](https://doi.org/10.1016/S0006-8993(87)80029-X).
59. Russell, W.M.S., and Burch, R.L. (1959). The Principles of Humane Experimental Technique. *eweb:789*. <https://repository.library.georgetown.edu/handle/10822/762203>.
60. Junca, P., Carcaud, J., Moulin, S., Garnery, L., and Sandoz, J.-C. (2014). Genotypic Influence on Aversive Conditioning in Honeybees, Using a Novel Thermal Reinforcement Procedure. *PLoS One* 9, e97333. <https://doi.org/10.1371/journal.pone.0097333>.
61. Percie du Sert, N., Hurst, V., Ahluwalia, A., Alam, S., Avey, M.T., Baker, M., Browne, W.J., Clark, A., Cuthill, I.C., Dirnagl, U., et al. (2020). The ARRIVE guidelines 2.0: Updated guidelines for reporting animal research. *J. Cerebr. Blood Flow Metabol.* 40, 1769–1777. <https://doi.org/10.1177/0271678X20943823>.
62. Junca, P., and Sandoz, J.-C. (2015). Heat Perception and Aversive Learning in Honey Bees: Putative Involvement of the Thermal/Chemical Sensor AmHsTRPA. *Front. Physiol.* 6, 316. <https://doi.org/10.3389/fphys.2015.00316>.
63. Koo, T.K., and Li, M.Y. (2016). A Guideline of Selecting and Reporting Intraclass Correlation Coefficients for Reliability Research. *J. Chiropr. Med.* 15, 155–163. <https://doi.org/10.1016/j.jcm.2016.02.012>.
64. Holm, S. (1979). A Simple Sequentially Rejective Multiple Test Procedure. *Scand. J. Stat.* 6, 65–70.

STAR★METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|----------------------------------------|-----------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------|
| Deposited data | | |
| Behavioral data | Figshare | https://doi.org/10.6084/m9.figshare.24534622.v1 |
| Code | Figshare | https://doi.org/10.6084/m9.figshare.24498304.v3 |
| Experimental models: Organisms/strains | | |
| <i>Bombus terrestris</i> bees | Biobest Group, Belgium | NA |
| Software and algorithms | | |
| R Studio | R Core Team, Cran-r-project, Vienna, Austria, version 2022.12.0 + 353 | NA |
| BORIS | Version 7.9.15; Italy | NA |

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Matilda Gibbons (matilda.gibbons@pennmedicine.upenn.edu).

Materials availability

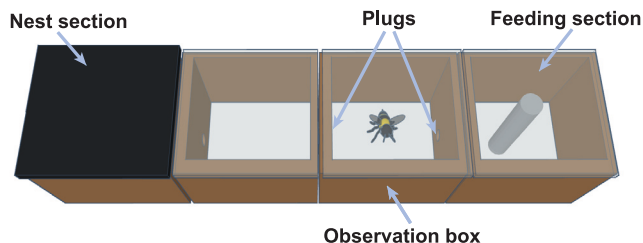
This study did not generate new unique materials.

Data and code availability

- Grooming duration data have been deposited at Figshare and are publicly available as of the date of publication. DOIs are listed in the [key resources table](#).
- All original code has been deposited at Figshare and is publicly available as of the date of publication. DOIs are listed in the [key resources table](#).
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

We used 82 adult bees from seven bumblebee colonies (standard hives from Biobest Group, Belgium). The bees were group-housed in ventilated wooden boxes (56 × 16 × 11 cm; see below figure). Each box comprised four sections, arranged linearly and connected by 1cm-diameter holes. At one end was the section containing the nest, which was covered with plywood. The section at the opposite end contained a 35 mL cylindrical feeder (74.5 × 31 mm), which dispensed Biogluc sugar solution *ad libitum* (Biobest group, Belgium). To access the food source, the bees had to cross the middle two sections. The middle section adjacent to the feeding section was the observation box during the testing period. The floor of both middle sections was covered with a thin layer of cat litter (Catsan Hygiene Plus, Mars Inc, USA) to absorb waste and debris. Each colony received 7g of pollen (Natupol Pollen, Koppert Biological Systems) every two days, and the laboratory was maintained at 23°C. We sexed each bee visually post-testing from the videos, based on the presence (in females) or absence (in males) of a black abdomen tip. There were 40 females and 18 males. The effect of sex on grooming behavior is discussed in the results. Bees from the same colony were pseudo-randomly assigned to experimental groups (pseudo-random to ensure there were bees in each experimental group from each colony).



Housing and testing apparatus

A ventilated wooden box (56 × 16 × 11 cm) with four sections. The nest section was covered with plywood. The feeding section contained a feeder with *ab libitum* food. The observation box was adjacent to the feeding section. Rubber plugs prevented the bee from leaving the observation box during the experiment.

METHOD DETAILS

Treatments

The United Kingdom does not regulate insect welfare in research. Nonetheless, we followed the 3Rs principles⁵⁹ in our experimental design and husbandry. In this vein, although some noxious stimulation is required to study self-protective behavior, we chose a temperature that, when brief, has no long-term effects on bees (65°C; based on⁶⁰). We also used a power analysis to estimate the minimum required sample size (estimated sample size = 80; alpha: 0.05; power: 80%). According to current best practice, we have followed the ARRIVE guidelines for reporting this research.⁶¹

For testing, we removed bees individually from the nest box by letting them walk onto metal forceps and placed them into a marking cage (Thorne, UK). A sponge in the marking cage was used to temporarily immobilize the bees to ensure precisely targeted noxious stimulation. A soldering iron (HAKKO FX-888D; Japan) was either heated to 65°C (noxious condition) or not heated (tactile condition), then touched onto the right or left antenna (counterbalanced across bees) for 5 s. We chose this method of noxious stimulation based on how stimulation of a honeybee's (*Apis mellifera*) antenna with a 65°C heat probe causes consistent sting extension reflexes⁶² (a defense reflex seen in response to noxious stimuli⁵²). Thirty bees were touched with the noxiously heated probe (noxiously stimulated; $N = 30$); 28 were touched with the control unheated probe (tactilely stimulated; $N = 28$); and 24 were put in the marking cage but not touched with a probe (control; $N = 24$). No bees were excluded from the analysis. We used an RST Soldering Iron Tip Thermometer 191 (YWBL-WH; China) to ensure the correct temperature of the soldering iron. After the treatment, bees were immediately placed in the observation box and filmed with an iPhone 8 (Apple; USA) for 25 min. We sealed the holes between boxes during the experiment, so bees were confined to the observation box (14 × 16 × 11 cm).

Behavioral analysis

Four treatment-blind coders recorded the self-grooming behavior displayed in the 25-min videos using BORIS behavioral analysis software (BORIS, version 7.9.15; Italy). Self-grooming was defined as 'the right or left front, middle, or hind leg moves over the left or right antenna either in one direction or in a repeated back and forth motion'. To measure inter-rater reliability, all four raters recorded grooming behavior for two bees (corresponding to two 25-min videos: one noxiously stimulated bee and one tactilely stimulated bee). Because the rating scale was continuous, we calculated the intraclass correlation coefficient. The correlation compared the total grooming duration of the right and left antenna across the four raters. The coefficient was 0.86, on a scale of 0–1, indicating a 'good' reliability.⁶³

QUANTIFICATION AND STATISTICAL ANALYSIS

We analyzed the data in R Studio (R Core Team, Cran-r-project, Vienna, Austria, version 2022.12.0 + 353), using general linear mixed effect models (GLMMs; packages: 'lme4' and 'car') and Wilcoxon tests. We checked model assumptions using histograms and 'Q-Q plots', and corrected for multiple testing using the Holm-Bonferroni correction.⁶⁴ We considered $p < 0.05$ significant and define N as number of bees. Sample size was determined by a power analysis (estimated sample size = 80; alpha: 0.05; power: 80%). Statistical details of the experiments can be found in the results section and in Figure 2.

To test for a difference between the grooming duration on the touched versus untouched antenna in noxiously stimulated and tactilely stimulated bees, we ran a GLMM. The response variable was the duration of antennal grooming for each antenna per bee. The fixed effects were stimulation type (noxious or tactile), whether the antenna was touched or untouched, the sex of the bee and their interaction. The random effect was the bee identity. We ran this model for the whole observation period (25 min), as well as individual time bins 0–1, 0–2, 0–3, 0–4 0–5, 6–10, 11–15, 16–20 and 21–25 min. We tested the individual time bins because some previous invertebrate studies have only detected self-grooming within the first few minutes after stimulation.^{17–19} The only time bin with a significant interaction effect (after applying the Holm-Bonferroni correction for multiple testing) was 0–2 min, so this is the only time bin we ran the other GLMM and Wilcoxon tests on (described below).

We used unpaired two-sample Wilcoxon tests (as our data did not meet the criteria for parametric analysis) to test the difference between the grooming durations on the touched or untouched antenna in the tactile and noxious treatment groups in the first 2 min after stimulation.

We ran another GLMM to test for a difference between grooming durations on either the touched or untouched antenna in the noxiously stimulated and tactilely stimulated bees, and the mean grooming duration for both antennae in bees in the control condition in the first 2 min. The response variable was either the duration of grooming on the touched antenna per bee or the duration of grooming on the untouched antenna per bee, or, for control bees, the mean grooming on one antenna was used, because neither antenna was touched in this condition. The fixed effects were the stimulation type (noxious, tactile, control) and the sex of the bee. The random effect was bee identity.