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Odour conditioning of positive affective states: Rats can learn to associate an odour with being tickled

Vincent Bombail1,*, Nathalie Jerôme1, Ho Lam1,2, Sacha Muszlak1, Simone L. Meddle2,3, Alistair B. Lawrence3,4, Birte L. Nielsen5

1 Neurobiology of Olfaction, INRA, Université Paris-Saclay, Jouy-en-Josas, France, 2 University of Edinburgh, Edinburgh, United Kingdom, 3 The Roslin Institute, The Royal (Dick) School of Veterinary Studies, Midlothian, United Kingdom, 4 Scotland’s Rural College, Edinburgh, United Kingdom, 5 UMR Modélisation Systémique Appliquée aux Ruminants, INRA, AgroParisTech, Université Paris-Saclay, Paris, France

* vincent.bombail@inra.fr

Abstract

Most associative learning tests in rodents use negative stimuli, such as electric shocks. We investigated if young rats can learn to associate the presence of an odour with the experience of being tickled (i.e. using an experimenter’s hand to mimic rough-and-tumble play), shown to elicit 50 kHz ultrasonic vocalisations (USVs), which are indicative of positive affect. Male, pair-housed Wistar rats (N = 24) were all exposed to two neutral odours (A and B) presented in a perforated container on alternate days in a test arena. Following 60s of exposure, the rats were either tickled on days when odour A (n = 8) or odour B (n = 8) was present, or never tickled (n = 8). When tickled, rats produced significantly more 50 kHz USVs compared to the days when not being tickled, and compared to control rats. The level of anticipatory 50 kHz USVs in the 60s prior to tickling did not differ significantly between the tickled and control rats. As a retrieval test following the odour conditioning, rats were exposed successively in the same arena to three odours: an unknown neutral odour, extract of fox faeces, and either odours A or B. Compared to controls, 50 kHz USVs of tickled rats increased when exposed to the odour they had previously experienced when tickled, indicating that these rats had learned to associate the odour with the positive experience of being tickled. In a test with free access for 5 min to both arms of a T-maze, each containing one of the odours, rats tickled with odour A spent more time in the arm with this odour. This work is the first to test in a fully balanced design whether rats can learn to associate an odour with tickling, and indicates that positive odour conditioning has potential to be used as an alternative to negative conditioning tests.

Introduction

Aversive conditioning, where a previously neutral stimulus or place becomes associated with an aversive experience, can be used to study memory and other brain functions in laboratory
rodents [1]. This paradigm is used in studies of learning, as odours can be associated with aversive states such as fear [2,3] and malaise [4,5]. The animals usually learn this association very quickly, making it a time-saving and efficient research method, which may be why only few attempts have been made to develop tests for this purpose using positive experiences. In studies where positive conditioning of odours has been applied, they involved pairing with psychostimulant drugs [6–8], alcohol [9] or a food source [10–12]. However, using feed as the unconditioned stimulus is not always feasible in practice, is likely to be associated with an increasing level of satiety, and psychoactive drugs alter mood and cognition [7].

We were therefore interested in finding an appropriate positive conditioning stimulus for use in an associative learning test for rats. This would ideally consist of a stimulus that was easy to use and which gave rise to the animal experiencing positive welfare (i.e. a positive affective state and not just absence of negative welfare; [13]). Despite the increasing interest in positive welfare indicators, the vast majority of animal welfare research has been and continues to be focused on more negative aspects [14]. One result of this is that there are few well-validated models of positive welfare in animals. One of the best candidates is the rat tickling model that was developed to mimic the effects of social play, a behaviour frequently displayed by young rats (see [15] for a recent review). In this model, the human hand is used to mimic the tactile stimulation experienced during social play in rats. The model has been validated partly through the measurement of ultrasonic vocalisations (USVs) that rats produce under different emotional states. During tickling, rats produce many more frequency modulated USVs in the range of 33–100 kHz (henceforth referred to as 50 kHz USVs); these are sometimes referred to as 'laughter' [16], and have been shown to indicate a positive emotional state [17]. Tickled rats show shorter latencies to approach the human hand than do controls, and express so-called optimistic biases when appraising environmental cues [18]. In addition, a number of pharmacological manipulations of rats using various psychotropes supports the notion that 50 kHz USVs are produced upon activation of the brain’s reward pathways [19,20]. Data thus support the interpretation that expressions of 50 kHz USVs indicate that tickling is a positive experience for the rat. However, USVs in the range of 22 kHz are emitted by rats under aversive situations [21–24]. USVs have therefore been suggested to be a useful tool for inferring affective states of the rats [25–28].

In this paper, we present the results of an experiment which was designed to condition rats to associate the presence of an odour with the positive experience of tickling. We hypothesised that if rats learned to make the odour-tickling association, they would i) emit anticipatory USVs when exposed to the odour prior to being tickled, ii) emit more 50 kHz USVs than control rats when exposed to the conditioned odour following exposure to an aversive odour, and iii) would spend more time in the arm of a T-maze containing their tickling odour.

Materials and methods

Animals and housing

Male Wistar rats (n = 24) were bred at the local animal facility at INRA, Jouy-en-Josas (permission N° A 78-322-5) and used as subjects for odorant conditioning and subsequent behavioural testing. The rats were weaned in groups of 6 at 21 days and housed in pairs at 4 weeks of age in standard laboratory rodent cages (42.5 cm × 26.6 cm × 18.5 cm made from transparent polycarbonate; Techniplast 1291H) on a 4-tier rack. The lighting schedule of the room was inverse 12D:12L, with lights coming on at 19:00 hours. The cages had a metal grid lid with a dentation in which commercial rat pellets (Diet M25, Special Diet Services, Witham, Essex, United Kingdom) were placed for ad libitum access. Water was supplied via a drinking bottle with a metal spout, inverted and placed alongside the feed. The floor of the cage was covered
by 2 cm of sawdust litter changed weekly, and wooden chew sticks (12 cm long) were supplied as enrichment. Individual rats were identified by marker pen lines on the tail. The rats were weighed once a week and, if needed, their tails were remarked.

**Odour conditioning schedule**

The rats were handled daily by the same person, and all handling, conditioning and testing took place during the dark period. Over the course of 5 days, the rats were gradually habituated to being put into a transport box (identical to the home cage, but with no water and feed available), and transported within an opaque black sack to the conditioning room, which was illuminated by red incandescent bulbs. The rats were also habituated to the conditioning arena—initially in pairs and subsequently individually. The arena consisted of a Plexiglas tank (LxWxH: 66 cm x 41 cm x 41 cm), bedded with sawdust (Fig 1A and 1B). At one end of the arena, a thin metal plate was fixed centrally at the bottom of the wall. During habituation, an empty stainless steel container (diameter 9.5 cm; height 3.7 cm; Grundtal IKEA) with a magnetic base and a screw-top lid with a perforated plastic inset was affixed vertically to the metal plate. This type of container was used to present the odour source to the rat during odour conditioning and subsequent testing.

In order to account for potential odour dependent effects, two different odours, previously found to be neutral to rats [29–31], were used as the conditioning odours: **odour A** (a 10% dilution of D-limonene; CAS no. 5989-27-5) and **odour B** (a 5% dilution of 1-hexanol; CAS no. 111-27-3); both diluted in mineral oil (CAS no. 8042-47-5). Different concentrations were used in order to ensure that the intensity of the two smells were as similar as possible, without being overpowering but strong enough to ensure a rapid detection by the rats. Using the same concentration of two different compounds does not per default ensure a similar intensity, and intensity of an odour is only vaguely related to the molecular weight and the vapour pressure of the compound [32]. Therefore, it is essential to assess experimental odour perception empirically, and a number of studies have shown the human nose is a detector equivalent to that of other species [33]. For each of the two odours used, we therefore presented five samples in decreasing dilutions to five of our colleagues, and chose the highest dilution (i.e. the weakest intensity) which was detectable by at least two people. All compounds were purchased from Sigma Aldrich (Saint-Quentin, Fallavier, France).

Each rat was allocated to one of three conditioning treatments, and all rats were placed in the arena on every conditioning day, where the conditioning treatment for each rat was applied: **A-tickled** rats were tickled when odour A was present in the container in the test arena; **B-tickled** rats were tickled when odour B was present in the container in the test arena; and **Control** rats were never tickled but still presented with either of the odours in the arena on alternate days (Fig 1C). When odour A was in the container, the B-tickled rats were treated as the Control rats, as were the A-tickled rats when odour B was in the container. Each pair of rats within a cage was randomly allocated to a treatment, whilst ensuring that rats housed on each tier of the rack received all three treatments. The order in which the rats were treated changed from one day to the next, starting with rats 1, 19, 13, and 7 on different days, respectively.

Conditioning began when the rats were 6 weeks of age. Only one of the two odours were used on each conditioning day to minimise the risk of cross-contamination and all testing was carried out in a well-ventilated room. Prior to the conditioning of each rat, a 2-ml sample of the odour dilution was transferred to a cotton pad in the container. All conditioning sessions where video recorded (Sony 12.0 mega pixels HDR-XR-500 Handycam) and the USVs registered using a freeware sound-recording programme (Audacity 2.1.3; www.audacityteam.org)
via a USV sensitive (10 to 160 kHz) microphone (M500-384, Pettersson Elektronik, Sweden) attached to the arena at an angle of 45° above the container (Fig 1A and 1B). A session began by moving the rat from its home cage in the transport box to the conditioning room and placing the rat in the arena 10 cm from and facing the container containing one of the two odours.

**Odour conditioning schedule:**

- **Type:** Anticipatory
- **Duration:** 60 s

| Session 1 | ... | Session 5 |
|-----------|-----|-----------|
| Day       |     |           |
| 1         |     |           |
| 2         |     |           |
| ...       |     |           |
| 9         |     |           |
| 10        |     |           |

| Odour      | Day 1 | Day 2 | Day 9 | Day 10 |
|------------|-------|-------|-------|--------|
| Control (n=8) | X     | X     | X     | X      |
| Tickled with A (n=8) | V     | X     | V     | X      |
| Tickled with B (n=8) | X     | V     | X     | V      |

**Fig 1. Odour conditioning schedule.** Screen-shots from the video recording of an odour conditioning session, showing a) the rat being tickled, and b) the position of the hand flat against the arena wall during the pauses between tickling periods. The container with the odour source can be seen in the left of the pictures. The pictures have a green hue as the procedure was carried out under red lighting; c) Diagram of the odour conditioning schedule over days and within treatment, with only the 1st (days 1 and 2) and 5th (days 9 and 10) session shown. Control rats were never tickled and A-tickled rats were tickled when odour A was present in the container, and B-tickled rats were tickled when odour B was present in the container in the test arena. When odour A was in the container, the B-tickled rats were treated as the Control rats, as was the A-tickled rats when odour B was in the container.
A or B. The rat was left there for 1 min, both to acclimatize from the move and to record anticipatory USVs upon exposure to the odour. This anticipatory period was kept relatively short to ensure that any tickling would occur soon after exposure to the tickling odour. The rat was subsequently either Tickled (A-tickled rats on days when odour A was present, B-tickled rats on days when odour B was present) or Not Tickled (Control rats on all days, A-tickled rats when odour B was present, and B-tickled rats when odours A was present; Fig 1C) according to the following procedures:

Tickled consisted of the handler using one hand wearing a knitted glove to touch, tickle, and play with the rat for 20 second periods interspersed with 20 second pauses. During the active periods, the handler mimicked the rough-and-tumble play seen in adolescent rats, with the hand tickling, chasing and pinning the rat, depending on its response (Fig 1A). After 20 seconds the hand was placed flat on the inside of the wall of the arena (Fig 1B). It rested here for 20 seconds after which another tickling period was carried out. This was repeated for a total duration of 140 s, allowing 4 periods of tickling interspersed with 3 periods of pauses. At the end of the tickling, the rat was moved in the transport box to a holding cage in a room separate from their home cage. The holding cage was identical to the home cage of the rat, placed in a similar 4-tier rack in the same position, and one holding cage was used for each pair of rats. The tickled rats were left in the holding cage for 3–7 hours (depending on the test order of the day) to prevent emotional contagion by USVs of the yet untested rats in the home cages.

Not Tickled consisted of the handler placing the hand wearing a knitted glove flat on the inside of the wall of the arena. It rested here for 20 seconds after which the hand was moved to the adjacent wall for 20 seconds. This was repeated for a total duration of 140 s, allowing 4 and 3 periods, respectively, with the hand resting on each wall, the latter being identical to the pauses when the rats were being Tickled. When the Not Tickled procedure finished, the rat was moved in the transport box back to its home cage.

In two pilot studies, we first tried to present the odour on the glove worn by the handler whilst tickling. However, this did not allow us to investigate anticipatory USVs before the hand was present in the arena. Also, in the pilot studies, the rats were moved back to their home cage after each conditioning session, and we were unsure if USVs emitted by tickled rats upon return would affect the yet untested rats via emotional contagion [34]. It has been found [35] that three tickling sessions sufficed to bring about a higher rate of 50 kHz USVs in tickled rats, and in a pilot study using only one odour, we found a significant difference in 50 kHz USVs emitted between tickled and control rats after four tickling days [36]. Nevertheless, as two odours were used alternately in the present experiment, we chose to carry out a total of ten conditioning days, alternating between odours A and B (Fig 1C). This resulted in all tickled rats being exposed to the Tickled procedure five times and the Not Tickled procedure five times, and all the rats were exposed to both odours for the same amount of time over the course of the ten days, including the Control rats.

For subsequent data analyses, the 50 kHz USVs for each rat were counted for the first (days 1 and 2) and the fifth (days 9 and 10) conditioning sessions, using the method described by [37]. Briefly, all USVs over 33kHz were defined as 50 kHz calls; these were not divided into subtypes (e.g. trills, step-calls) but counted with individual calls being separated by at least 0.048 s. These counts were divided into 50 kHz USVs emitted during the four tickling periods (4 x 20s) and the three pauses (3 x 20s), as well as the 1-min habituation period prior to tickling to detect any differences in anticipatory USVs. This grouping of 50 kHz USVs was also done when the Not Tickled procedure was applied (Fig 1C). All counts were converted into USVs/min. Occurrences of 22 kHz USVs were rare, emitted in only 5 conditioning sessions out of the 96 analysed, with the majority being during the first session; these are therefore not included in the data.
The behaviour of the rats was logged from the video recordings of the 5th session (days 9 and 10). This was done for the three 20s pauses only, as the hand did not move and the behaviour of the rats at this time was therefore comparable within and among all rats and across treatments and procedures. The recorded behaviour consisted of hand seeking behaviour (the rat rears to sniff the motionless hand, see Fig 1A, or is facing and focusing on the hand), play jumping (jumping while running), sniffing the air, exploring the odour container, freezing (immobility, often sudden, with ears raised and eyes open) and other behaviour (locomotion, digging the litter, and self-grooming).

**Behavioural tests of conditioning**

On the two days following the last conditioning session, two behavioural tests were carried out to investigate the effects of the conditioning:

In the Triple Odour test, the rat was placed in the same arena as used for the conditioning. The tests consisted of 30-sec periods with no odour container present in the arena, interspersed with three 1-min periods, where a container was positioned containing the following odours in said order: 1) a neutral odour unknown to the rat (Novel odour; a 5% suspension of p-anisaldehyde, CAS no. 123-11-5, in mineral oil), which was assumed to have no aversive or attractive properties for the rats; 2) an extract in mineral oil of fox faeces (Fox odour; faecal pellets originating from several male foxes and soaked in mineral oil for 24h at 70˚C, with extract diluted 1:6); this odour was expected to induce a level of fear in the rats, and 3) the Tickling odour with which the Tickled rats had been conditioned, and with half of the Control rats being exposed to odour A and the other half to odour B. As the test arena was large and had no lid, we assessed the 30s inter-odour period to be sufficient to disperse the previous odour, whilst maintaining the interest of the rats at each odour placement. The order of the three odours were chosen so as to measure the response of the rats to first an unknown, but neutral odour, then an unknown but fear-inducing odour, followed by the known conditioning odour. We hypothesised that if the Tickled rats had learned to associate their tickling odour with a positive experience, more 50 kHz USVs would be emitted by the Tickled rats compared to the Control group when exposed to the conditioning odour, the latter having been exposed to the odour for the same amount of time during conditioning but without being tickled. The tests were video recorded and the latencies to explore the odour container and the amount of freezing displayed by the rats during exposure to the three different odours was scored. The USVs were registered in the same way as for the conditioning sessions.

In the T-maze test, the rat was placed without prior habituation at one end of a large T-maze arena, which consisted of a rectangular open space (77 cm x 51 cm) with two accessible arms (WxL: 19 cm x 25 cm) extending from each side at one end of the rectangle, forming a broad T-shape. A ventilator fitted centrally at the other end of the rectangle extracted air from the T-maze, ensuring a simultaneous airflow from both arms. No litter was used, and a perforated metal tea-ball was placed in a pre-drilled hole at the end wall of each arm of the maze. The two tea-balls each contained a cotton pad imbibed with 2 ml of either odour A or B, with one odour in each arm, alternating between arms in a balanced way for each rat being tested. The test was video recorded from above. The rat was left in the arena for 5 min, and was free to explore both arms and the central arena.

All housing and experimental procedures for this experiment were carried out in accordance with European legislation on the protection of animals used for scientific purposes [38], and had been approved by the local ethics committee (Comité d’Ethique en Expérimentation Animale INRA IdF-Jouy-en-Josas/ AgroParisTech (Comethea); permission #16–17). At the
end of the experiment, the animals were returned to the main cohort for use in other studies, as encouraged by [38] when mild procedures have been used.

**Statistical data analysis**

Data were analysed in MiniTab (ver. 17.1) using General Linear Models (GLM) followed by post-hoc Bonferroni comparisons of significant effects. For the USVs emitted during conditioning, data from the four tickling periods were analysed fitting odour, treatment, and session with interaction. When relevant, Pearson’s correlations were calculated. Anticipatory USVs were analysed for session 5 only, fitting odour, treatment and their interaction. Behaviour during pauses, expressed as percentage of time spent on each behaviour, was analysed by the non-parametric Kruskal-Wallis test, as data were not normally distributed. For the Triple Odour test, differences in USVs emitted as the test progressed as well as freezing behaviour were compared between treatment groups using GLM, with levels prior to the first odour presentation fitted as a covariate to adjust for individual differences. Data from the T-maze test were analysed using GLM across odours as well as among tickling treatments, adjusting for rat within treatment. Results are given as means ± standard errors, unless otherwise stated.

**Results**

All tickled rats emitted significantly more 50 kHz USVs during the sessions with the Tickled procedure than during the sessions when Not Tickled ($F_{2,89} = 31.3; P < 0.001$), with the latter not differing in magnitude from that of the never tickled Control rats (Fig 2A). This was evident already during the very first tickling session, but with significantly more 50 kHz USVs emitted during the 5th compared to the 1st tickling session (233 vs 83 (±8.4) USVs/min; $P < 0.001$), and these were significantly correlated (Pearson’s $r = 0.53; P = 0.036$), indicating that response level of 50 kHz USVs to tickling is a characteristic of the individual rat. No differences in USV frequency were found between rats tickled with Odours A and B, respectively (235 vs 230 (se = 11.7) USVs/min; $F_{1,28} = 0.8; P = 0.383$). On days when the tickled rats were Not Tickled (i.e. A-tickled rats when odour B was present and vice versa), the 50 kHz USVs emitted per minute by the 5th tickling session was higher than that found in the 1st session and did not differ significantly from the level observed during tickling in the 1st session, indicating a degree of place association had developed (Fig 2A), whereas the control rats did not show a significant increase with session.

Rats also emitted 50 kHz USVs during the pauses between tickling periods. When rats were tickled, although numerically greater, the USVs during pauses in the 1st session did not differ significantly from those observed during the tickling (108 vs 83 (±11.1) USVs/min; $P = 0.390$), whereas by the 5th session, significantly more 50 kHz USVs were emitted during tickling than during the pauses (151 vs 233 (±11.1) USVs/min for pauses and tickling periods, respectively; $P < 0.001$; Fig 2C and 2D). For the Not Tickled rats, including Controls, the levels of 50 kHz USVs were similar between tickling periods and pauses because no actual tickling took place. When rats were tickled, 50 kHz USVs emitted during pauses and during tickling were correlated for the 1st ($r = 0.76; P = 0.001$) but not the 5th session ($r = 0.02; P = 0.955$). The latter was caused by the expected increase in USV emissions during actual tickling. When data excluding the tickled rats were analysed, the correlation was significant (Pearson’s $r = 0.78; P < 0.001$), reflecting a greater variability among than within individual rats.

Medians of the behaviour during pauses in the 5th session are shown in Table 1, with tickled rats showing significantly more hand seeking behaviour and play jumping, with consequently less time spent in general locomotion, than when Not Tickled or compared to Control rats. Using the same data, a comparison between the odours across treatments found a small, but significant difference in the percentage of time sniffing the air (Medians: odour A: 27.5%,
During the pauses, no significant differences between odours A and B were found across rat treatments in USVs emitted (86 ±13.7 vs 83 ±10.2 USVs/min, respectively; P = 0.856), nor during the anticipatory periods (57 ±6.8 vs 56 ±8.1 USVs/min, respectively; P = 0.808).

Even after five tickling sessions, rats did not emit more anticipatory USVs when exposed to their tickling odour, compared to when exposed to their non-tickling odour and compared to
the control group ($F_{2,42} = 0.33; P = 0.718$; Fig 2B). Fig 3 shows the frequency of 50 kHz USVs emitted by the rats as a function of their anticipatory 50 kHz USVs. The higher level of USVs during tickling is clearly visible, but with no clear correlation with anticipatory USVs for the tickled rats. However, for the sessions without tickling (Not Tickled and Control rats), the 50 kHz USVs emitted show a positive relationship with the 50 kHz USVs emitted during the pre-session (anticipatory) minute ($R^2 = 53.1\%; T = 4.99; P < 0.001$; Fig 3), supporting previously reported findings that rats may be characterised according to their level of vocalisation [39].

Table 1. Behaviours observed during pauses between tickling.

| Behaviour (median % of time spent) | Control rats | Not Tickled rats | Tickled rats | P ≤ |
|-----------------------------------|--------------|------------------|-------------|-----|
| Sniffing the air                  | 34.2         | 30.2             | 28.3        | 0.198|
| Locomotion                        | 29.2         | 32.5             | 12.5        | 0.001|
| Hand seeking behaviour            | 12.5         | 18.5             | 35.0        | 0.001|
| Exploring odour container         | 3.3          | 3.3              | 1.67        | 0.839|
| Play jumping                      | 0.0          | 0.8              | 10.8        | 0.001|
| Freezing                          | 0.0          | 1.7              | 4.2         | 0.168|
| Digging the litter                | 0.8          | 2.5              | 0.0         | 0.057|
| Self-grooming                     | 0.8          | 0.0              | 0.0         | 0.060|

Median percentage of time spent in different behaviours by rats during pauses between tickling periods in the 5th conditioning session. Control rats were never tickled, and data for Not Tickled rats are from the session with the non-conditioning odour, when the Tickled rats were subjected to the Not Tickled procedure. Within rows, a median in bold font differ significantly from the other medians.

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Fig 3. Data plot of 50 kHz ultrasonic vocalisations (USVs) per minute during the 5th session in the periods when tickling occurred for the Tickled rats plotted against the USVs per minute during the 60-s period (anticipatory) prior to tickling. Each data point is an individual rat, with Control rats plotted for both Odours A and B. The regression equation, where data from Tickled rats have been excluded, is $Y = 1 + 0.78X$ ($R^2 = 58\%$; $P < 0.001$).

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In the Triple Odour test, rats approached and sniffed the odour container within a few seconds, independent of the odour (overall median = 3 s; Q3-Q1 = 5.5). USVs emitted during the Triple Odour test are shown in Fig 4A. During the first minute, where no odour was present, the frequencies of 50 kHz USVs (overall mean: 69 ±8.0 USVs/min) were no different from those seen during the anticipatory period during odour conditioning (see Fig 2B). These data were used as a covariate in the analysis to adjust for individual differences in base level USV emissions among the rats. Over the course of the Triple Odour test, USV frequency decreased gradually, but a significant increase in 50 kHz USVs from the preceding pause was found for the tickled rats when exposed to their tickling odour (increase: 20±5.7, 32±5.7 and –4±5.8 USVs/min for A-tickled, B-tickled, and Control rats, respectively; F<sub>2,20</sub> = 10.3; P < 0.001), with the increase being significantly different for both A-tickled (P = 0.024) and B-tickled rats (P < 0.001) from that of the controls. The rats thus increase their 50 kHz USV frequency when their conditioning odour was presented indicating that the rats had learned to associate an odour with the positive experience of tickling.

Freezing was scored during the Triple Odour test, as exposure to fox odour was expected to induce more freezing as an indicator of fear. However, as shown in Fig 4B, freezing did not increase when fox odour was in the arena, and only one rat emitted 22kHz USVs during the test, starting when the fox odour was introduced. The Tickled rats showed the same low level of freezing throughout the test, independent of odour present. The Control rats, however, showed a significant increase in freezing when one of the conditioned odours were present, and this was mainly due to greatly elevated levels of freezing in the Control rats (n = 4) exposed to odour A (F<sub>3,19</sub> = 9.0; P = 0.001; Fig 4B).

Behaviour during the T-maze test showed that overall, more time was spent in the arm with odour A (60.6 ±4.45 vs 44.8±4.45 s for odours A and B, respectively; F<sub>1,23</sub> = 6.6; P = 0.017). This was significantly different from time spent in the arm with odour B for the A-tickled rats (F<sub>1,14</sub> = 5.0; P = 0.041; Fig 5A and 5B) with a similar tendency for Control rats (P = 0.088). The proportion of time spent in the arm with odour A was significantly different from chance (0.5) only for the rats tickled with odour A (0.60; T = 2.51; P = 0.041).

**General discussion**

Using an appetitive conditioning method, we aimed for rats to learn to associate the presence of an odour with the positive experience of tickling. Our first hypothesis was that rats, which had learned to make the odour-tickling association would emit more anticipatory USVs when exposed to the odour prior to being tickled. This was not the case, as no differences in anticipatory USVs were found between the treatment groups. Indeed, our use of the term anticipatory can be questioned, given the findings. Rats will emit 50 kHz calls in anticipation of access to a running wheel [40]. Others have found that individual rats vocalize more in a chamber associated with play than in a habituated control chamber [41], indicative of anticipation of positive experiences. The absence of similar anticipatory vocalisations in the present experiment would indicate that, with respect to our first hypothesis, the conditioning paradigm was not successful. However, the levels of USVs emitted during the (anticipatory) pre-session minute appeared to predict the overall level of vocalisation for individual rats when these were not tickled, suggesting that rats can be categorised according to their USV frequency independent of any tickling occurring. This is in accordance with [39,42], who divergently selected rats based on their 50 kHz vocalisations. It has previously been found that tickling-induced 50 kHz ultrasonic vocalisations are individually stable and can predict behaviour in tests of anxiety and depression in rats [43,44]. Our finding that USVs produced when not being tickled show large inter-individual differences, but little intra-individual variation, is complementary to these results.
Fig 4. USVs and freezing behaviour during the triple odour test. a) Mean number of 50 kHz ultrasonic vocalisations (USVs; ± s.e.) per minute, and b) mean duration (s; ± s.e.) of freezing behaviour during the Triple Odour test for A-tickled odour conditioning of positive affective states.
The second hypothesis was that more USVs would be emitted by the tickled than by control rats when exposed to the conditioned odour following exposure to an aversive odour. We found an increase in USVs produced by the tickled rats when their tickling odour was placed in the arena. Given that 50 kHz USVs are indicative of positive affect [25], this would indicate that the tickled rats had learned to associate the odour with a positive experience. Ideally, we would have tested the conditioned rats with both odours, but the small number of animals made this statistically inappropriate. However, the increase in 50 kHz USVs by the A-tickled and B-tickled rats when exposed to their conditioned odour in the Triple Odour test was not simply because the odour was known compared to the two previous odours, as the Control rats showed no such increase in USV production. It was noted that exposure to the fox odour did not provoke freezing behaviour in the rats and only elicited 22 kHz USVs in a single rat, indicating that this odour was less aversive than anticipated.

We also expected tickled rats to spend more time in the arm of a T-maze containing their tickling odour. This was found only for rats tickled in the presence of odour A, and across treatments rats spent longer in the arm containing odour A than the arm containing odour B. As we did not want the rats to have experienced the odours used before the conditioning started, we did not test the relative attractiveness of the odours before carrying out the T-maze test, which might have strengthened the interpretability of these test results. The overall increase in air sniffing during conditioning when odour B was in the arena corresponds to the findings from the Triple Odour test, where odour B appeared to have a stronger effect than odour A (see Fig 4A). Control rats also showed more freezing when exposed to odour A in the Triple Odour test. Although freezing is often considered an indication of fear, the behaviour is but a display of increased alertness, and the interpretation is context specific. Exposure to oestrous odours can elicit freezing in rats [30,45] and this is enhanced by sexual experience [31]. It may be that odour A induced freezing in control rats because they find it more interesting, as shown by their un-conditioned preference in the T-maze test. The two odours were chosen as being neutral to rats [29], and we struggle to explain why they affect the behaviour of the rats differently. One, speculative possibility is a potential sedative effect of inhaling limonene resulting in decreased locomotor activity, which has been found in mice [46,47], but no differences in activity of the rats were found among treatments during odour conditioning. Even neutral odours may differ in their response eliciting properties, even if they are intrinsically neither aversive nor attractive to the rats.

As mentioned in the introduction, aversive conditioning is a widely used technique in learning studies of memory and other brain functions in laboratory rodents [1,4,5]. Pairing aversive stimuli such as electric shocks, with a neutral stimulus or situation usually give rise to associations learned within a few sessions [2]. In contrast, the application of appetitive conditioning regimes often require more pairing sessions to become effective. Although our positive conditioning was successful, as shown by the response of the rats to their conditioning odour in the Triple Odour test, this did not appear to be a very strong association, as no increase in anticipatory vocalisation was seen as conditioning progressed, nor a very convincing preference for the tickling odour in the T-maze test. Others have also struggled to demonstrate a link between increased 50 kHz USVs and reward-related stimuli: [37] conditioned rats to associate a tone with a food reward, and measured the expression of reward anticipation as increases in USVs. However, when the rats were food-deprived, they showed only behavioural but not
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**Figure a)**

**Time (s) spent in arm**

- **Odour A**
- **Odour B**

Control: 60 sec
A-tickled: 70 sec
B-tickled: 60 sec

**Figure b)**

Image of a T-maze setup with labeled sections A and B.
vocal anticipation and when sated, the reward cue continued to elicit 50 kHz USVs despite being devalued by pre-feeding. These findings may, in part, have been due to the large inter-individual variability among rats, giving rise to different types of responders [48]. It is evident from these and the present results that appetitive conditioning is likely to be more complex and less effective than most aversive conditioning.

One protocol of rat tickling has been described in detail by [49]. This consists of 15s pauses in between 15s tickling bouts with 4 to 5 dorsal contacts and pins each time. Systematic tickling procedures allow for comparisons across experiments. However, we did not standardise the tickling method used in the present experiment, over and above the fixed alternating periods of tickling and pauses. This was a conscious choice on our part, as we had previously found a large individual variation in the response of the rats to tickling. Our experience indicated that this variation was reduced if the rats were tickled and played with whilst allowing the hand to react to the behavioural responses of the individual rat. In addition, as tickling is a playful experience, it should be varied and unpredictable to the rats. Although the lack of standardisation prevented us from comparing behaviour of the rats during the active tickling period, i.e. as the behaviour of the experimenter varied slightly across rats and across sessions, we were able to use the hand-seeking behaviour during the pauses to assess the likability of tickling for each rat. Tickled rats showed more hand seeking behaviour and play jumping with simultaneously more 50 kHz USVs emitted during the pauses between tickling, indicating that the tickling lead to a positive affective state.

In conclusion, rats learned to associate an odour with the positive experience of being tickled, as they increased their 50 kHz USVs when exposed to this odour in a test situation without tickling, compared to control rats that had been exposed to the same odour for the same amount of time without being tickled. However, no increase was seen in anticipatory USVs when exposed to the conditioning odour prior to being tickled, and only one of the conditioning odours gave rise to a preference by the tickled rats in a T-maze test. These findings indicate that rats can learn to associate an odour with the positive experience of tickling, and positive odour conditioning may thus have potential to be developed further with a view to replacing negative odour conditioning tests. However, different odours may differ in their efficacy, and appetitive (positive) conditioning is clearly more difficult and slower to induce than aversive (negative) conditioning. This experiment is but the first step to develop behavioural tests for use in the laboratory employing positive affective states in the conditioning paradigm.

Supporting information

S1 Dataset. Data used in the statistical analyses. Data on 50 kHz ultrasonic vocalisations (USVs) in sessions 1 and 5 and in the Triple Odour test, as well as behaviour data from session 5, Triple Odour test and the T-Maze test.

(XLSX)

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Author Contributions

**Conceptualization:** Vincent Bombail, Ho Lam, Alistair B. Lawrence, Birte L. Nielsen.

**Data curation:** Nathalie Jérome, Ho Lam, Birte L. Nielsen.

**Formal analysis:** Vincent Bombail, Birte L. Nielsen.

**Funding acquisition:** Vincent Bombail, Simone L. Meddle, Alistair B. Lawrence.

**Investigation:** Vincent Bombail, Nathalie Jérome, Alistair B. Lawrence.

**Methodology:** Vincent Bombail, Nathalie Jérome, Ho Lam, Sacha Muszlak, Simone L. Meddle, Alistair B. Lawrence, Birte L. Nielsen.

**Project administration:** Vincent Bombail, Alistair B. Lawrence, Birte L. Nielsen.

**Resources:** Simone L. Meddle.

**Software:** Vincent Bombail, Nathalie Jérome, Sacha Muszlak.

**Supervision:** Birte L. Nielsen.

**Validation:** Sacha Muszlak, Alistair B. Lawrence.

**Visualization:** Sacha Muszlak.

**Writing – original draft:** Birte L. Nielsen.

**Writing – review & editing:** Vincent Bombail, Nathalie Jérome, Ho Lam, Sacha Muszlak, Simone L. Meddle, Alistair B. Lawrence, Birte L. Nielsen.

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