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induced by learning and illness in mice

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### ABSTRACT

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A male-specific major urinary protein named darcin is attractive to female mice, stimulates a learned attraction to volatile components of a male's urinary odour (Roberts et al., 2010, BMC Biol 8:75) and induces spatial learning (Roberts et al., 2012, Science 338:1462-5). In the present work we show that darcin also induces a learned attraction for a previously neutral olfactory stimulus (the odorant isoamyl acetate). However, the attractive properties of darcin may change as a function of female physiological state. For example, during the period of lactation female mice display aggressive behaviour against intruders, which is enhanced when confronted with adult males. Therefore, the endocrinological status of the females radically changes the behavioural response. The situation at puberty is somewhat similar: prepubertal females avoid adult male chemical signals, whereas post-pubertal females show attraction for these same signals. In addition, we report another situation in which the presence of darcin is not attractive to adult female mice. Urine of males parasitized by the intestinal nematode Aspiculuris tetraptera shows no attractive value for female mice, despite apparently normal presence of darcin. In this case, the loss of the attractive value is not due to physiological changes in the receptor females but is likely to be due to the presence of unknown signals of infection whose detection overrides the attraction normally induced by darcin.

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- KEY WORDS: vomeronasal, olfactory, sexual attraction, learning, maternal aggression,
- 38 puberty, illness cues.

Pheromones were originally defined as "substances which are secreted to the outside by an individual and received by a second individual of the same species, in which they release a specific reaction, for example, a definite behavior or a developmental process" (Karlson & Luscher, 1959). Although this definition has been very useful for more than 50 years (Wyatt, 2009), the response to pheromones (at least in mammals) may vary depending on a number of factors, including the previous experience or the hormonal status of the receiver (Wyatt, 2010). Moreover, the behavioural response elicited by pheromones may also be observed as learned responses to stimuli previously associated with the pheromones, so that these stimuli acquire pheromone-like properties (Moncho-Bogani, Lanuza, Hernández, Novejarque, & Martínez-García, 2002). Here, we review relevant examples in mice, in which the attractive value of male sexual pheromones is transferred to a neutral stimulus or changes either as a function of the hormonal condition of the receiver females or the health status of scent donor males. In addition, we include data from two experiments giving support to the ideas presented in this review.

## Pheromone-induced olfactory learning

Odours are easily associated with either positive or negative experiences, becoming secondary attractive or aversive stimuli that strongly influence behavioural responses in many mammalian species, including humans (Herz & Cupchik, 2005). In rodents, olfactory stimuli play a key role in many aspects of socio-sexual behaviours (Brennan & Kendrick, 2006). The experience with social chemical signals influences later responses. For example, chemical signals present in urine of male mice, detected through the vomeronasal system, have reinforcing properties able to induce appetitive associative learning (Martínez-Ricós, Agustín-Pavón, Lanuza, & Martínez-García,

- 65 2007, 2008; Moncho-Bogani et al., 2002; Ramm, Cheetham, & Hurst, 2008), in such a
- way that other volatiles present in urine may become secondary attractive odorants
- 67 (Martínez-García et al., 2009). More recently, a male-specific urinary protein named
- darcin, has been shown to be able to induce this kind of olfactory learning (Roberts et
- al., 2010) and also spatial learning (Roberts, Davidson, McLean, Beynon, & Hurst,
- 70 2012). We hypothesized that darcin would be able to induce a secondary attraction by
- association with a neutral odorant (not present in urine). To test this possibility, we
- 72 performed the following experiment.
- 73 EXPERIMENT 1: Inducing attraction for a neutral odorant by association with darcin
- 74 Material and Methods
- 75 Subjects
- For the present study, 15 adult female mice (12-16 weeks) of the CD1 outbred strain
- 77 were used (Janvier Labs., Le Genest-Saint-Isle, Saint-Berthevin Cedex, France).
- 78 Treatment of the animals employed in the experiments reported in this paper complied
- with the European Union Council Directive of June 3<sup>rd</sup>, 2010 (6106/1/10 REV1),
- 80 according to which procedures were approved by the Committee of Ethics on Animal
- 81 Experimentation of the University of Valencia (protocol number A1283764105250).
- 82 Procedures also adhered to the ASAB/ABS Guidelines for the Use of Animals in
- 83 Research.
- The females were sexually naïve and had never been exposed to chemical signals from
- sexually mature males. To achieve this, pregnant females were housed in a clean room
- without males, in standard macrolon transparent cages with a wire lid (21.5 x 46.5 x
- 87 14.5 cm, ref. 1000, Panlab, Barcelona, Spain) filled with soft wood bedding (Souralit
- 88 S.L., ref. 3000, Barcelona, Spain), provided with nesting material (shredded paper) and
- 89 enriched with cardboard tubes. The room was maintained at 22-24 °C, 60–80% RH and

a 12:12 h light:dark cycle, with lights on at 0800 hours). Food (Teklad Global 14% Protein Rodent Maintenance Diet, Harlan, ref: 2014) and water were available ad libitum. Nineteen days after delivery (early before puberty), pups were sexed and males were removed. Their female siblings were brought to a clean room in complete absence of adult male chemical signals, where they were kept in groups of 5-6 per cage (the stock housing conditions in the experimental room were the same as described above for the pregnant females). Food and water were available ad libitum except during the preference tests (five-minute long) and the training sessions (15-minute long). Welfare assessment took place during cage cleaning, and included non-invasive indicators. In the neonates, skin colour, activity and presence of the milk spot were observed; at weaning and in the adult, general appearance, size, coat condition, posture, gait, activity levels, interaction with the environment and clinical signs were observed (Wells et al., 2006). After weaning, animals were only manipulated for cage cleaning once a week. Since general appearance and size were evaluated as normal, no further care was necessary. Mice were handle following the standard practice of picking them up by gently holding the base of the tail and helping them onto the handler's arm, avoiding to hold them in the air. All procedures involved in this study were non-invasive behavioural tests. At the end of the experiments, animals were euthanized with an intraperitoneal overdose of sodium pentobarbital (92 mg/kg), as indicated in the approved protocol (cited above). The male siblings were either used for anatomical studies (protocols approved by the Committee of Ethics on Animal Experimentation of the University of Valencia, under the same reference number A1283764105250; published elsewhere, Otero-García et al., 2014) or euthanized as described above. Stimuli

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114 We chose two odorants that have been used frequently as olfactory stimuli in the 115 literature: isoamyl acetate (e.g., Angely & Coppola, 2010; Panreac, Barcelona, Spain) 116 and citralva (e.g., Martínez-Ricós et al., 2007; geranonitrile, 3,7-dimethyl-2,6-117 octadiene-1-nitrile, kindly provided by International Flavours and Fragrances, Ventos, 118 Barcelona, Spain). Both odorants were diluted 1:1000 in phosphate buffer (0.01 M) with 119 0.01% Triton X-100. In a pilot test of olfactory preference we proved that isoamyl 120 acetate and citralva were investigated equally. 121 Preference tests 122 Animals were habituated to handling and to the test cage over 3 days, 10 min per day, 123 between 15:00 and 20:00 hours. Preference tests were performed in 25 x 50 x 30 cm 124 cages. A 4 x 4 cm piece of filter paper impregnated with 5 µl of one of the stimulus 125 odorants was presented in each opposing side of the cage. These impregnated papers 126 were fixed to the bottom of the cage with a metallic cover, leaving exposed a circular 127 area (diameter = 3.5 cm) that allowed direct nasal contact with the paper but prevented 128 the animals to gnaw or remove it. 129 For this olfactory preference test (citralva vs isoamyl acetate) females were released in 130 the centre of the cage, the experimenter left the room, and the behaviour was videotaped 131 for 5 min. The time that the animal spent in the circular area covered by the paper was 132 measured by tracking animals automatically using the video analyser software Smart 133 2.5 (Panlab, Barcelona, Spain; see Fig. 1A). Since we observed that the animals lost 134 interest in the olfactory stimuli at the end of the test, we restricted the analysis of the 135 data to the first four minutes. 136 Following the pre-training preference test, the females were run in a second test in 137 which the isoamyl acetate-scented paper was also impregnated with 8 µl of recombinant 138 darcin (r-darcin, diluted 1.1 µg/µl, Roberts et al., 2010). Over the next four days, four

training sessions (one per day) were performed in which a piece of paper (not fixed) scented with 5 µl of isoamyl acetate and 8 µl of r-darcin was presented to the females during 15 min/day in the centre of a different cage (29 x 15 x 29.8 cm), and thus in a different context to that of the preference tests. Finally, a post-training olfactory preference test was performed (citralva vs isoamyl acetate) identical to the pre-training test described above. The location (left or right) of the citralva and isoamyl acetatescented papers was decided randomly at the beginning of the experiment and kept fixed for all the animals and tests. After finishing this experiment, we wanted to discard the possibility that repeated presentations of the odorant, by themselves, induced preference for this familiar stimulus (against the other odorant, which is less familiar). To test this idea, we performed a second (control) experiment (n = 12, 16 weeks of age, Janvier Labs., Saint-Berthevin Cedex, France) identical to the previous one, except for the absence of darcin in the preference test and training sessions. Since females of this control experiment were not going to be exposed to male-derived chemosignals, they were acquired from Janvier as young adults, without preventing their prepubertal exposure to male odours. Housing and care conditions were the same as described above. Statistical analysis The time spent in the area occupied by the scented paper was analysed with a repeatedmeasures ANOVA with the TEST (control, 1st odour preference test; darcin preference test; 2<sup>nd</sup> odour preference test) and SIDE (citralva vs. isoamyl acetate) as intra-subjects factors. The normality of the data was previously confirmed with a Kolmogorov-Smirnov test with Lillieford's correction. Analyses were performed with the SPSS 15.0 software package.

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Results

164 The results of the repeated measures ANOVA for the time spent within each of the circular areas (SIDE) during the four TESTS (Control; 1st odour preference test; darcin 165 preference test; 2<sup>nd</sup> odour preference test; Figure 1B) revealed significant main effects 166 167 of SIDE ( $F_{1.14} = 7.87$ , p = 0.014), and non-significant main effect of TEST or the SIDE 168 x TEST interaction (F < 1, p > 0.6 in both cases). The analysis of the simple effects of 169 the factor SIDE in each test showed that both areas were equally investigated in the control (clean versus clean, F < 1, p > 0.4) and the 1<sup>st</sup> odour preference test (citralva vs. 170 171 isoamyl acetate,  $F_{1,14} = 1.18$ , p = 0.16). By contrast, the females spent significantly more time in the area occupied by the paper scented with isoamyl acetate plus darcin 172  $(F_{1,14} = 5.07, p = 0.032)$ . Finally, in the  $2^{nd}$  (post-training) olfactory preference test 173 174 (citralva vs. isoamyl acetate), females again spent more time in the area occupied by the 175 isoamyl acetate-scented paper ( $F_{1.14} = 5.77$ , p = 0.031). 176 In the control group, in which the same procedure was used but no darcin was present, 177 the results of the repeated measures ANOVA for the time spent within each of the 178 circular areas (SIDE) during the four TESTS (Figure 1B) showed no significant main 179 effects of SIDE ( $F_{1,11} < 1$ , p = 0.51), TEST ( $F_{3,9} < 1$ , p = 0.71), or their interaction ( $F_{3,9}$ 180 = 2.06, p = 0.17). The analysis of the simple effects of the factor SIDE in each test showed that both areas were equally investigated in all cases (clean versus clean:  $F_{1.11}$  = 181 182 3.23, p = 0.1; first odour preference test:  $F_{1.11} = 1.02$ , p = 0.33; second odour preference 183 test:  $F_{1,11} < 1$ , p = 0.99; third odour preference test:  $F_{1,11} = 1.37$ , p = 0.26). 184 Discussion 185 The results of the present experiments show that a neutral odorant, such as isoamyl 186 acetate, which is not significantly preferred by female mice, becomes a preferred 187 olfactory stimulus by presenting it together with the sexual pheromone darcin. The 188 repeated presentation of the odorant, as shown by the control experiment, did not alter

189 the original lack of preference between the two olfactory stimuli used in the present 190 tests. 191 Darcin is a male-specific non-volatile urinary protein (MUP20, MW 18893 Da) 192 previously shown to induce in females a learned olfactory preference for the particular 193 pattern of urinary volatiles displayed by an individual mouse (Roberts et al., 2010). In 194 addition, darcin can also induce spatial learning (female mice also remember the 195 location where it was presented in a test cage, Roberts et al., 2012). Regarding this, we 196 should keep in mind that spatial learning is also likely to take place in the present 197 experiments, since we ran a five-minute preference test in which darcin was present 198 following the first citralva vs. isoamyl acetate test. However, some relevant differences 199 between the present experiments and those reported by Roberts et al. (2012) suggest a 200 weaker role of spatial learning in the present case. Firstly, we used a test cage with no 201 internal spatial cues, with both sides of the cage being equal. Secondly, our test was five 202 minute long (the training session in Roberts et al., 2012, was 10 minutes long). Thirdly, 203 we used 8 µl of r-darcin, while 50 µl were used in Roberts et al. (2012), at 204 approximately the same concentration. And finally, in the present experiment the 205 females were later exposed to isoamyl acetate scented papers impregnated with darcin 206 daily for 15 min over the next four days, with these sessions taking place in a very 207 different context. This provided abundant possibilities for the formation of odour-208 pheromone associations, whereas the opportunities for spatial learning were 209 comparatively more reduced. In any case, the possible role of spatial learning cannot be 210 discarded, and future experiments should confirm the induction of odour-pheromone 211 learning suggested by the present results. 212 Previous work has shown that isoamyl acetate can be used as a conditioned stimulus in 213 an aversive learning task, associating it with lithium chloride (Kay & Nyby, 1992), and

214 therefore this olfactory stimulus can be conditioned to be either aversive or attractive. 215 We hypothesize that in the present case of associative learning the conditioned stimulus 216 (isoamyl acetate) is detected by the olfactory system and the unconditioned stimulus 217 (darcin) is detected by the vomeronasal system (since the animals need direct contact 218 with the stimulus to show either innate attraction or learned associations, Roberts et al., 219 2010). Olfactory and vomeronasal information are known to converge in several nuclei 220 within the corticomedial amygdala (Cádiz-Moretti, Martínez-García, & Lanuza, 2013; 221 Kang, Baum, & Cherry, 2009, 2011; Pro-Sistiaga et al., 2007), where learning may take 222 place. In addition, further intramygdaloid projections would allow the participation of 223 the nuclei of the associative (basolateral) amygdala, as suggested by functional data 224 obtained with the immediate early gene c-Fos (Moncho-Bogani, Martínez-García, 225 Novejarque, & Lanuza, 2005). A different pheromone that has been shown to induce 226 olfactory learning is the rabbit mammary pheromone 2-methylbut-2-enal (2MB2) 227 (Coureaud et al., 2006), although in this case the 2MB2 is likely detected by the main 228 olfactory system, as indicated by functional studies using the Fos protein as neural 229 activity marker (Charra, Datiche, Gigot, Schaal, & Coureaud, 2013). 230 The induction of a learned preference for airborne urinary stimuli should take place in 231 natural conditions when female mice explore the urine marks that males use to advertise 232 their territory (Hurst & Beynon, 2004). The urine of males is enriched in several volatile 233 molecules, such as farnesenes, 2-sec-butyl 4,5 dihydrothiazole, and 3,4 dehydro-exo-234 brevicomin, which have been shown to be also detected by the vomeronasal organ 235 (Leinders-Zufall et al., 2000) and maybe possess pheromonal activity on their own (see, 236 for a review, Dulac & Torello, 2003). For example, the mixture of alpha and beta 237 farnesenes was shown to be attractive to sexually naïve female mice but only when used 238 in very high concentrations, while having no effect when used at a concentration that

was double that of normal dominant male urine (Jemiolo, Xie, & Novotny, 1991). By contrast, farnesenes were preferred even at low concentrations by sexually experienced animals (Jemiolo et al., 1991). Although the previous chemosensory experience of the animals in these experiments is unknown, the effects of sexual experience clearly indicate a role for learning. Similar remarks can be made in other cases of putatively identified pheromonal stimuli, such as (methylthio)methanethiol (MTMT, an attractive semiochemical present only in the urine of male mice, Lin, Zhang, Block, & Katz, 2005) and androstenone, a pheromone that facilitates expression of both attraction to the male and a receptive mating stance in estrous female pigs (Dorries, Adkins-Regan, & Halpern, 1997). In both studies the female subjects had previous sexual experience (in the case of the female pigs most of them were multiparous). In the light of the results presented here, the pheromonal role of these semiochemicals should be reevaluated at least using sexually naïve (if not chemosensory naïve) animals to understand the requirement for learning. In the same vein, the human steroid androstenone (5α-androst-16-en-3-one) has been proposed to function as a human sex chemosignal (see, for a review, Havlicek, Murray, Saxton, & Roberts, 2010). However, the hedonic value of androstenone was recently evaluated as a function of sexual experience (Knaapila et al., 2012), the odour being rated as unpleasant by women who reported never having experienced sexual intercourse, and as less unpleasant by those who reported being sexually experienced. Since humans do not have a functional vomeronasal organ (Meredith, 2001), in this case learning is likely to be mediated by the association of the olfactory cue with other kinds of rewarding stimuli related to sexual activity. The phenomenon of pheromone-induced olfactory learning raises the question of whether a substance that gains its role as chemical signal by a learned association should be considered a pheromone, since it does not elicit a fixed (stereotyped) response

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(as required by the original definition of Karlson & Luscher, 1953) before learning takes place. However, in the case of the male-specific airborne urinary substances this olfactory-vomeronasal association would necessarily occur every time the female interacts with males of male urine marks. Moreover, females will encounter different olfactory-vomeronasal associations with each male, so that the learned response would be specific to that particular signature (Ramm et al., 2008). Under natural conditions, females are probably able to detect male-specific cues at a distance using the main olfactory system. Once the female locates the male (or his urine marks) the input through the vomeronasal organ (requiring direct contact with the source) would allow further information about this particular individual to be processed. Several of the male-specific urinary volatiles are also detected by the vomeronasal organ (Leinders-Zufall et al., 2000), but it is currently unknown why the detection of volatile signals by the vomeronasal organ requires direct contact with the stimulus, as indicated by both behavioural and electrophysiological evidence (Luo, Fee, & Katz, 2003; Moncho-Bogani et al., 2002; Ramm et al., 2008).

# Turning attraction into aggression: male-specific vomeronasal cues elicit maternal aggression

As stated above, the attraction that females display for male urine is innate. However, during lactation females show aggressive responses towards male (to a lesser extent also towards female) intruders, to protect their pups (maternal aggression, Rosenson & Asheroff, 1975). Maternal aggression towards intruders is observed in the first two weeks after delivery, and disappears gradually onwards (Lonstein & Gammie, 2002). Maternal aggression is low towards castrated males, and vomeronasal organ removal in females prior to mating or after parturition eliminates later maternal aggression in mice

(Bean & Wysocki, 1989). Therefore, for the female to show maternal aggression the vomeronasal detection of testosterone-dependent chemical stimuli from males is required. This raises the question of whether some of the major urinary proteins, whose synthesis is testosterone-dependent and have been shown to be the vomeronasal stimuli mediating aggression between males (Chamero et al., 2007), are also the vomeronasal stimuli that elicit maternal aggression in lactating females. Preliminary data in our laboratory suggest that this is indeed the case (Martín-Sánchez et al., 2013), and that the attractive pheromone darcin is also able to induce maternal aggression. Therefore, the hormonal status of the lactating females induces changes in the neural structures processing darcin (probably not in the vomeronasal organ, although direct evidence of this is lacking). These changes may take place in the amygdalo-hypothalamic circuits involved in the aggressive response (Nelson & Trainor, 2007), but experimental evidence of this hypothesis is needed. Notably, lactating females are not aggressive towards familiar mates (Lonstein & Gammie, 2002), and therefore the response also depends on the learned identity of the individual male. Individual recognition in mice is mediated by the pattern of major urinary proteins (Hurst et al., 2001), and therefore the detection of the pattern of major urinary proteins corresponding to the familiar male should inhibit the aggressive response. Social recognition involves changes of gene expression (in particular oxytocin, vasopressin and steroid hormone receptors) in the amygdala and the hypothalamus (Clipperton-Allen et al., 2012), giving support to the hypothesis of the control of the aggressive responses by amygdalo-hypothalamic circuits (Bosch & Neumann, 2012).

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Turning aversion into attraction: changes at puberty inducing attraction to male pheromones

Although the sexual attraction that female mice display for the male sexual pheromone darcin does not require learning (Roberts et al, 2010), it seems to appear with puberty, since pre-pubertal females display an aversive response to chemical signals from unfamiliar adult males (Drickamer, 1989; Mucignat-Caretta, Caretta, & Baldini, 1998). The biological bases of this change are unknown, although clearly the gonadal steroids underlying the pubertal changes should play a relevant role. In fact, the vomeronasal organ, likely involved in the detection of protein sex pheromones (such as darcin, Roberts et al., 2010; and exocrine gland-secreting peptide 1, Haga et al., 2010), as well as most of the neural centres of the vomeronasal system are sexually dimorphic (Segovia & Guillamon, 1993). This suggests a role for sexual steroids in development and differentiation of the vomeronasal system. Moreover, gonadal steroids also regulate the expression of some vomeronasal receptors (Alekseyenko, Baum, & Cherry, 2006), raising the possibility that sex steroids induce the expression of particular (currently unknown) receptors for male pheromones. In fact, there is some evidence of estradiol effects on the induction of c-Fos in the vomeronasal organ of female mice by malesoiled bedding (Halem, Cherry, & Baum, 1999). In addition, the presence of estradiol receptors is very important in the secondary vomeronasal centres, namely the medial amygdala, the posteromedial cortical amygdala and the posteromedial bed nucleus of the stria terminalis (Mitra et al., 2003). Therefore, the pubertal changes underlying the induction of the attraction of female mice for male pheromones may also take place in the amygdaloid circuits processing vomeronasal information.

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From attraction to aversion: illness-derived chemicals and avoidance of conspecifics

339 sexual pheromones and neurobiological foundation of this phenomenon. Occasionally 340 we observed that viral infections or parasitosis in male mice used as donors of urine or 341 bedding resulted in a lack of attractiveness of the urine (or soiled bedding) for females. 342 This fits previous studies reporting avoidance of males by females when males were 343 infected with viruses or parasites (e.g. Penn et al. 1998; Kavaliers et al., 2005). To 344 check the response of female mice to the urine of infected males experimentally, we 345 performed preference tests (Experiment 2) using urine from males parasitized by the 346 nematode Aspiculuris tetraptera against (healthy) female urine. In addition, since darcin 347 appears to be the both necessary and sufficient to make male urine attractive to females, 348 we tested whether the urine of parasitized males contains darcin. 349 EXPERIMENT 2. Behavioural response of female mice to the urine of infected males 350 Material and Methods 351 Subjects and stimuli 352 For this experiment, 32 adult female mice (12-16 weeks) of CD1 strain were used 353 (Janvier, Le Genest-Saint-Isle, Saint-Berthevin Cedex, France). As for experiment 1, 354 females were sexually naïve and had never been exposed to chemical signals from 355 sexually mature males. Females were housed in groups of 5-6 animals, with the housing 356 conditions (cages, bedding and food), manipulation and welfare assessment being the 357 same as those reported in Experiment 1. 358 Urine from healthy adult CD-1 male and female mice was purchased from Janvier and 359 kept frozen in aliquots until used. Urine from a small colony (n = 4) of male mice (also 360 purchased from Janvier) naturally infected with the intestinal nematode Aspiculuris 361 tetraptera was collected as described by Kurien, Everds & Scofield (2004). Briefly, 362 animals were gently held by the scruff of the neck over a petri dish, from where urine

During the last years, our research has been focused on the attractive properties of

was pipetted. Since the infection occurred naturally, the animals can probably experience this level of parasitism in the wild. The presence of these parasites in male mice was detected with the routine sentinel vigilance system. The infected mice were sacrificed except for four animals that were kept for 5 days. During this time we collected their urine once a day. At the end of this 5-day period the four infected animals were also sacrificed. The infected mice showed no external signs of infection, and general appearance, size, coat condition, posture, gait, activity levels, interaction with the environment and clinical signs appeared normal. Infected male mice were housed in pairs in standard macrolon cages with a wire lid (22.5 x 22.5 x 14.5 cm, ref. 500, Panlab, Barcelona, Spain). The rest of housing conditions were the same as described in Experiment 1. To ensure the homogeneity of the stimulus across behavioural tests, urine from different males was mixed and stored in frozen aliquots of 50 μl. Infection with the parasite was assessed through the presence of eggs in faecal pellets and confirmed post-mortem by checking for adult worms in the colon. Preference tests The test cage and habituation procedure were as described in experiment 1. Preference tests were performed in which the female mice had to choose between two urine stimuli located in opposite sides of the cage. To do so, 10 µl of the stimulus urine were pipetted on one of the tips of a rectangular piece of filter paper (2 x 6 cm) that was attached to the wall so that the urine spots were 8 cm above the floor. The females were able to have direct nasal contact with the stimuli by standing on the hind legs. Following a control test, with PBS 0.1M on both sides of the cage, the olfactory preference test (male vs female urine) was performed and recorded for five minutes as described for Experiment 1. In one group of animals (randomly assigned, n = 16), urine of healthy males was presented on one side and female urine on the other side. In the second group

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388 of females (randomly assigned, n = 16), urine of infected males was presented against 389 female urine. 390 Statistical analysis 391 The time spent in a semicircular area (of a radius of 3.85 cm) around the filter paper was 392 analysed with a two-way repeated-measures ANOVA with TEST (control, preference 393 test) and STIMULUS (male vs. female urine) as intra-subject factors, and GROUP 394 (urine from healthy or infected males) as an inter-subject factor. Significant interactions 395 were further analysed by multiple pair-wise comparisons with Bonferroni corrections. 396 The normality of the data was previously confirmed with a Kolmogorov-Smirnov test 397 with Lillieford's correction. Analyses were performed with the SPSS 15.0 software 398 package. 399 *Electrophoresis of urinary* proteins 400 The pattern of bands corresponding to urinary proteins, separated according to their 401 mass, was visualised using sodium dodecyl sulphate–polyacrylamide gel electrophoresis 402 (SDS-PAGE). Urine was diluted 1:1 with 2x denaturalization buffer (20mM Tris pH 403 8.0, 5% SDS, 10% mercaptoethanol, 2mM EDTA and 0.05% bromophenol blue) in a 404 capped Eppendorf tube, vortexed to mix, boiled for 5 min and then centrifuged for 5 405 min at 10000 rpm. The samples were then allowed to cool before sample loading. Using 406 1x denaturalisation buffer samples of male urine were brought to a final 1:6 dilution, 407 whereas female urine samples were not further diluted (final dilution 1:2). This allowed 408 direct comparison of male and female urine protein species, in spite of the difference in 409 total urinary protein content between sexes. Using a PhastGel system (General 410 Electrics), electrophoresis was run under reducing conditions at a constant 200 V on a 411 20% polyacrylamide gel (PhastGel Homogeneous – 20, GE). Low range molecular

weight markers (Sigmamarker low range, M3913, St. Louis, MO, USA) were used for

413 comparison. Following electrophoresis, protein bands were visualised using Phast Gel

Blue (0.1%) solution and differentiated in a solution of methanol: acetic acid: distilled

- 415  $H_2O$  (30:10:60 v/v/v).
- 416 Results

- The results of the repeated measures ANOVA of time spent within the areas
- surrounding the stimulus during the tests (Figure 2A) showed non-significant main
- 419 effects of factors TEST ( $F_{1.30} = 1.9$ , p = 0.17), STIMULUS ( $F_{1.30} < 1$ , p = 0.6) or
- 420 GROUP ( $F_{1,30} < 1$ , p = 0.9), and non-significant interactions between each pair of these
- 421 factors (STIMULUS x TEST:  $F_{1,30} = 1.14$ , p = 0.29; TEST x GROUP:  $F_{1,30} = 2.43$ , p = 0.29
- 422 0.12; STIMULUS x GROUP:  $F_{1.30} < 1$ , p = 0.4). However, there was a significant triple
- interaction (STIMULUS x TEST x GROUP:  $F_{1,30} = 7.45$ , p = 0.01). Pos-hoc analysis of
- 424 this triple interaction (Figure 2A) showed that in both groups the two stimulus areas
- were investigated equally in the control condition, when saline buffer was present on
- both sides of the cage (healthy male group, p > 0.7; infected male group, p > 0.9). By
- contrast, females presented with urine of healthy males showed a clear preference for
- 428 this stimulus over female urine (p = 0.023); females presented with urine of infected
- males spent more time next to the female urine, although the time spent next to each
- 430 stimulus was not significantly different (p = 0.27).
- To check for the presence of the sexual pheromone darcin in the urine of male mice
- infected with the nematode Aspiculuris tetraptera, we performed a SDS-PAGE
- electrophoresis to compare proteins in healthy male urine, urine of infected males, urine
- of castrated (non-infected) males and female urine (Figure 2B). The results showed that
- 435 the pattern of protein bands in the urine of infected males was similar to that observed
- in the urine of healthy males, with a clearly visible band of higher mobility than other
- major urinary proteins that corresponds to darcin (Armstrong et al., 2005). By contrast,

this band was not present in the urine from castrated males or in female urine,

confirming previous results (Armstrong, Robertson, Cheetham, Hurst, & Beynon, 2005;

Cheetham, Smith, Armstrong, Beynon, & Hurst, 2009).

Discussion

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It has been shown previously that female mice are able to discriminate between infected and non-infected males and show a preference towards healthy males (see Kavaliers, Choleris, & Pfaff, 2005). The results of the present experiment suggest that females not only choose non-infected males against infected males, but also that the preference for male urine over female urine is lost if urine comes from parasitized males. Since it is known that the attractive properties of male urine depend on the presence of a malespecific urinary protein named darcin (Roberts et al., 2010), which acts as a sexual pheromone, we tested whether the expression of this protein may have been lost in the infected males. The results show that darcin is present in the urine of parasitized males apparently at normal levels, though further analyses would be required to confirm whether there are quantitative differences. Although we cannot discard a small reduction in the expression of this protein, this is unlikely to explain the total lack of preference for male urine versus female urine that did not contain darcin (at least at a level that could be detected by electrophoresis). Changes in the amino-acidic sequence or conformation of the protein are very unlikely. Therefore, we can hypothesize that an infection cue exists in the urine of the parasitized animals, and that detection of this is able to override the attractive value of darcin. An alternative hypothesis, though, is that the cue of infection is volatile and can be detected at a distance; this detection may inhibit the vomeronasal pumping necessary to deliver high molecular weight molecules, such darcin, to the vomeronasal organ (Meredith, Marques, O'Connell, & Stern, 1980;

463 detected by females. 464 Whether volatile or involatile (or both), the identity of the putative cue(s) of infection is 465 unknown, as it is its nature as olfactory of vomeronasal stimulus. It has recently been 466 demonstrated that the vomeronasal organ of mice expresses a different type of 467 chemosensory receptor named FPR (formyl peptide receptors, Liberles et al., 2009; 468 Riviere, Challet, Fluegge, Spehr, & Rodriguez, 2009). This type of receptor, in addition 469 to formylated peptides produced by bacteria, detects ligands related with the immune 470 system (such the antimicrobial peptide CRAMP, lipoxin A4, or uPAR, see Riviere et

al., 2009), and therefore are good candidates for sensing of urinary infection cues,

although experimental evidence for this possibility is currently lacking.

Wysocki, Wellington, & Beauchamp, 1980). In this case, darcin would simply not be

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### GENERAL CONCLUSSIONS

475 Although the attractive response that adult female mice show towards the male-specific 476 pheromone darcin is innate, here we review four examples in which either the 477 behavioural response can be induced by non-pheromonal stimuli or the presence of the 478 pheromone does not induce the normal attractive response, due a non-receptive 479 hormonal status (pre-pubertal or lactating females) or to the putative presence of 480 infection cues in the male urine. 481 The induction of learned attraction by a pheromone is an interesting model to study 482 learning and memory mechanisms in an ethologically relevant context. In addition, 483 changes in response to darcin induced at puberty and during lactation provide 484 unexplored models of neural plasticity with clear and robust behavioural correlates, 485 which may be helpful to understand the neural circuits that induce both attractive and 486 aggressive responses.

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- 494 Alekseyenko, O. V., Baum, M. J. & Cherry, J. A. 2006. Sex and gonadal steroid
- 495 modulation of pheromone receptor gene expression in the mouse vomeronasal organ.
- 496 *Neuroscience*, **140**, 1349-1357. doi: 10.1016/j.neuroscience.2006.03.001.
- 497 Angely, C. J. & Coppola, D. M. 2010. How does long-term odor deprivation affect the
- olfactory capacity of adult mice? *Behavioral and Brain Functions*, **6**, 26-9081-6-26.
- 499 doi: 10.1186/1744-9081-6-26.
- 500 Armstrong, S. D., Robertson, D. H., Cheetham, S. A., Hurst, J. L. & Beynon, R. J.
- 501 2005. Structural and functional differences in isoforms of mouse major urinary proteins:
- a male-specific protein that preferentially binds a male pheromone. *The Biochemical*
- 503 Journal, **391**, 343-350.
- Bean, N. J. & Wysocki, C. J. 1989. Vomeronasal organ removal and female mouse
- aggression: the role of experience. *Physiology & Behavior*, **45**, 875-882.
- Bosch, O.J. & Neumann, I.D. 2012. Both oxytocin and vasopressin are mediators of
- maternal care and aggression in rodents: from central release to sites of action.
- 508 *Hormones and Behavior*, **61**, 293-303. doi: 10.1016/j.yhbeh.2011.11.002.
- Brennan, P. A. & Kendrick, K. M. 2006. Mammalian social odours: attraction and
- 510 individual recognition. *Philosophical Transactions of the Royal Society. Series B*,
- 511 Biological Sciences, **361**, 2061-2078. doi: 10.1098/rstb.2006.1931.
- 512 Cadiz-Moretti, B., Martinez, G. F. & Lanuza, E. 2013. Neural substrate to associate
- odorants and pheromones: Convergence of projections from the main and accessory
- olfactory bulbs in mice. In: *Chemical Signals in Vertebrates 12* (Ed. by M. L. East & M.
- 515 Dehnhard), pp. 3-16. New York: Springer Science.
- 516 Chamero, P., Marton, T. F., Logan, D. W., Flanagan, K., Cruz, J. R., Saghatelian,
- 517 A., Cravatt, B. F. & Stowers, L. 2007. Identification of protein pheromones that
- 518 promote aggressive behaviour. *Nature*, **450**, 899-902. doi: 10.1038/nature05997.

- Charra, R., Datiche, F., Gigot, V., Schaal, B., & Coureaud, G. 2013. Pheromone-
- 520 induced odor learning modifies Fos expression in the newborn rabbit brain. *Behavioral*
- 521 Brain Research, 237, 129-140. doi: 10.1016/j.bbr.2012.09.017.
- 522 Cheetham, S. A., Smith, A. L., Armstrong, S. D., Beynon, R. J. & Hurst, J. L. 2009.
- 523 Limited variation in the major urinary proteins of laboratory mice. *Physiology &*
- 524 *Behavior*, **96**, 253-261. doi: 10.1016/j.physbeh.2008.10.005.
- 525 Clipperton-Allen, A.E., Lee, A.W., Reyes, A., Devidze, N., Phan, A., Pfaff, D.W. &
- 526 Choleris, E. 2012. Oxytocin, vasopressin and estrogen receptor gene expression in
- relation to social recognition in female mice. *Physiology and Behavior*, **105**, 915-924.
- 528 doi: 10.1016/j.physbeh.2011.10.025.
- 529 Coureaud, G., Moncomble, A.S., Montigny, D., Dewas, M., Perrier, G., & Schaal,
- **B.** 2006. A pheromone that rapidly promotes learning in the newborn. *Current Biology*,
- **16**, 1956-1961.
- Dorries, K. M., Adkins-Regan, E. & Halpern, B. P. 1997. Sensitivity and behavioral
- responses to the pheromone androstenone are not mediated by the vomeronasal organ in
- domestic pigs. Brain, Behavior and Evolution, 49, 53-62.
- Drickamer, L. C. 1989. Odor preference of wild stock female house mice (Mus
- domesticus) tested at three ages using urine and other cues from conspecific males and
- females. *Journal of Chemical Ecology*, **15**, 1971-1987.
- 538 **Dulac, C. & Torello, A. T.** 2003. Molecular detection of pheromone signals in
- mammals: from genes to behaviour. *Nature Reviews Neuroscience*, **4**, 551-562.
- Haga, S., Hattori, T., Sato, T., Sato, K., Matsuda, S., Kobayakawa, R., Sakano, H.,
- Yoshihara, Y., Kikusui, T. & Touhara, K. 2010. The male mouse pheromone ESP1
- 542 enhances female sexual receptive behaviour through a specific vomeronasal receptor.
- 543 *Nature*, **466**, 118-122. doi: 10.1038/nature09142.
- Halem, H. A., Cherry, J. A. & Baum, M. J. 1999. Vomeronasal neuroepithelium and
- forebrain Fos responses to male pheromones in male and female mice. *Journal of*
- 546 *Neurobiology*, **39**, 249-263.

- Havlicek, J., Murray, A. K., Saxton, T. K. & Roberts, S. C. 2010. Current issues in
- 548 the study of androstenes in human chemosignaling. *Vitamins and Hormones*, **83**, 47-81.
- 549 doi: 10.1016/S0083-6729(10)83003-1.
- Herz, R. S. & Cupchik, G. C. 1995. The emotional distinctiveness of odor-evoked
- 551 memories. *Chemical Senses*, **20**, 517-528.
- Honda, K., Negoro, H., Dyball, R. E. J., Higuchi, T. & Takano, S. 1990. The
- osmoreceptor complex in the rat: Evidence for interactions between the supraoptic and
- other diencephalic nuclei. *The Journal of Physiology*, **431**, 225-241.
- Hurst, J.L., Payne, C.E., Nevison, C.M., Marie, A.D., Humphries, R.E., Robertson,
- 556 **D.H., Cavaggioni, A. & Beynon, R.J.** 2001. Individual recognition in mice mediated
- by major urinary proteins. *Nature*, **414**, 631-634.
- Hurst, J. L. & Beynon, R. J. 2004. Scent wars: the chemobiology of competitive
- signalling in mice. *BioEssays*, **26**, 1288-1298.
- Jemiolo, B., Xie, T. M. & Novotny, M. 1991. Socio-sexual olfactory preference in
- female mice: attractiveness of synthetic chemosignals. *Physiology & Behavior*, **50**,
- 562 1119-1122.
- Kang, N., Baum, M. J. & Cherry, J. A. 2011. Different profiles of main and accessory
- olfactory bulb mitral/tufted cell projections revealed in mice using an anterograde tracer
- and a whole-mount, flattened cortex preparation. *Chemical Senses*, **36**, 251-260. doi:
- 566 10.1093/chemse/bjq120; 10.1093/chemse/bjq120.
- Kang, N., Baum, M. J. & Cherry, J. A. 2009. A direct main olfactory bulb projection
- to the 'vomeronasal' amygdala in female mice selectively responds to volatile
- 569 pheromones from males. The European Journal of Neuroscience, 29, 624-634. doi:
- 570 10.1111/j.1460-9568.2009.06638.x.
- Karlson, P. & Luscher, M. 1959. Pheromones': a new term for a class of biologically
- 572 active substances. *Nature*, **183**, 55-56.

- Kavaliers, M., Choleris, E. & Pfaff, D. W. 2005. Genes, odours and the recognition of
- parasitized individuals by rodents. *Trends in Parasitology*, **21**, 423-429. doi:
- 575 10.1016/j.pt.2005.07.008.
- Kay, E. & Nyby, J. 1992. LiCl aversive conditioning has transitory effects on
- 577 pheromonal responsiveness in male house mice (Mus domesticus). Physiology &
- 578 *Behavior*, **52**, 105-113.
- Knaapila, A., Tuorila, H., Vuoksimaa, E., Keskitalo-Vuokko, K., Rose, R. J.,
- **Kaprio, J. & Silventoinen, K.** 2012. Pleasantness of the odor of androstenone as a
- 581 function of sexual intercourse experience in women and men. Archives of Sexual
- 582 *Behavior*, **41**, 1403-1408. doi: 10.1007/s10508-011-9804-7.
- Kurien, B.T., Everds, N.E. & Scofield, R.H. 2004. Experimental animal urine
- collection: a review. *Laboratory Animals*, **38**, 333-361.
- Leinders-Zufall, T., Lane, A. P., Puche, A. C., Ma, W., Novotny, M. V., Shipley, M.
- **T. & Zufall, F.** 2000. Ultrasensitive pheromone detection by mammalian vomeronasal
- 587 neurons. *Nature*, **405**, 792-796.
- Liberles, S. D., Horowitz, L. F., Kuang, D., Contos, J. J., Wilson, K. L., Siltberg-
- Liberles, J., Liberles, D. A. & Buck, L. B. 2009. Formyl peptide receptors are
- candidate chemosensory receptors in the vomeronasal organ. *Proceedings of the*
- 591 National Academy of Sciences USA, **106**, 9842-9847. doi: 10.1073/pnas.0904464106.
- Lonstein, J. S. & Gammie, S. C. 2002. Sensory, hormonal, and neural control of
- maternal aggression in laboratory rodents. Neuroscience and Biobehavioral Reviews,
- **26**, 869-888.
- Luo, M., Fee, M. S. & Katz, L. C. 2003. Encoding pheromonal signals in the
- accessory olfactory bulb of behaving mice. *Science*, **299**, 1196-1201. doi:
- 597 10.1126/science.1082133.
- Martin-Sanchez, A., Hernandez-Martinez, A., McLean, L., Beynon, R.J., Hurst,
- 599 J.L., Lanuza, E., Martinez-Garcia, F. 2013. When males become the enemy: maternal

- aggression is induced by the attractive male sexual pheromone darcin in mice. 5th
- 601 Parental Brain Conference. Póster. Regensburg, Germany.
- Martinez-Garcia, F., Martinez-Ricos, J., Agustin-Pavon, C., Martinez-Hernandez,
- **J., Novejarque, A. & Lanuza, E.** 2009. Refining the dual olfactory hypothesis:
- pheromone reward and odour experience. Behavioural Brain Research, 200, 277-286.
- 605 doi: 10.1016/j.bbr.2008.10.002.
- Martinez-Ricos, J., Agustin-Pavon, C., Lanuza, E. & Martinez-Garcia, F. 2007.
- Intraspecific communication through chemical signals in female mice: reinforcing
- properties of involatile male sexual pheromones. *Chemical Senses*, **32**, 139-148.
- Martinez-Ricos, J., Agustin-Pavon, C., Lanuza, E. & Martinez-Garcia, F. 2008.
- Role of the vomeronasal system in intersexual attraction in female mice. *Neuroscience*,
- 611 **153**, 383-395. doi: 10.1016/j.neuroscience.2008.02.002.
- Meredith, M. 2001. Human vomeronasal organ function: a critical review of best and
- 613 worst cases. *Chemical Senses*, **26**, 433-445.
- Meredith, M., Marques, D. M., O'Connell, R. O. & Stern, F. L. 1980. Vomeronasal
- pump: significance for male hamster sexual behavior. *Science*, **207**, 1224-1226.
- 616 Mitra, S. W., Hoskin, E., Yudkovitz, J., Pear, L., Wilkinson, H. A., Hayashi, S.,
- Pfaff, D. W., Ogawa, S., Rohrer, S. P., Schaeffer, J. M., McEwen, B. S. & Alves, S.
- **E.** 2003. Immunolocalization of estrogen receptor beta in the mouse brain: comparison
- with estrogen receptor alpha. *Endocrinology*, **144**, 2055-2067.
- Moncho-Bogani, J., Martinez-Garcia, F., Novejarque, A. & Lanuza, E. 2005.
- Attraction to sexual pheromones and associated odorants in female mice involves
- activation of the reward system and basolateral amygdala. The European Journal of
- 623 Neuroscience, **21**, 2186-2198.
- Moncho-Bogani, J., Lanuza, E., Hernandez, A., Novejarque, A. & Martinez-
- 625 **Garcia, F.** 2002. Attractive properties of sexual pheromones in mice. Innate or learned?
- 626 *Physiology & Behavior*, **77**, 167-176.

- Mucignat-Caretta, C., Caretta, A. & Baldini, E. 1998. Protein-bound male urinary
- 628 pheromones: differential responses according to age and gender. Chemical Senses, 23,
- 629 67-70.
- Nelson, R. J. & Trainor, B. C. 2007. Neural mechanisms of aggression. *Nature*
- 631 Reviews Neuroscience, **8**, 536-546. doi: 10.1038/nrn2174.
- Otero-Garcia, M., Martin-Sánchez, A., Fortes-Marco, L., Martínez-Ricós, J.,
- 633 Agustín-Pavón, C., Lanuza, E. & Martínez-García, F. 2014. Extending the socio-
- 634 sexual brain: arginine-vasopressin immunoreactive circuits in the telencephalon of mice.
- 635 *Brain Structure and Function* (in press). Doi: 10.1007/s00429-013-0553-3.
- Penn, D., Schneider, G., White, K., Slev, P. & Potts, W. 2010. Influenza infection
- 637 neutralizes the attractiveness of male odour to female mice (*Mus musculus*). *Ethology*,
- 638 **104**, 685-694. doi: 10.1111/j.1439-0310.1998.tb00102.x.
- 639 Pro-Sistiaga, P., Mohedano-Moriano, A., Ubeda-Banon, I., Del Mar Arroyo-
- Jimenez, M., Marcos, P., Artacho-Perula, E., Crespo, C., Insausti, R. & Martinez-
- Marcos, A. 2007. Convergence of olfactory and vomeronasal projections in the rat
- basal telencephalon. *The Journal of Comparative Neurology*, **504**, 346-362. doi:
- 643 10.1002/cne.21455.
- Ramm, S. A., Cheetham, S. A. & Hurst, J. L. 2008. Encoding choosiness: female
- attraction requires prior physical contact with individual male scents in mice.
- 646 Proceedings of the Royal Society, Series B, Biological Sciences, 275, 1727-1735. doi:
- 647 10.1098/rspb.2008.0302; 10.1098/rspb.2008.0302.
- Riviere, S., Challet, L., Fluegge, D., Spehr, M. & Rodriguez, I. 2009. Formyl peptide
- receptor-like proteins are a novel family of vomeronasal chemosensors. *Nature*, **459**,
- 650 574-577. doi: 10.1038/nature08029.
- Roberts, S. A., Simpson, D. M., Armstrong, S. D., Davidson, A. J., Robertson, D.
- H., McLean, L., Beynon, R. J. & Hurst, J. L. 2010. Darcin: a male pheromone that
- stimulates female memory and sexual attraction to an individual male's odour. BMC
- 654 *Biology*, **8**, 75-7007-8-75. doi: 10.1186/1741-7007-8-75.

- Roberts, S. A., Davidson, A. J., McLean, L., Beynon, R. J. & Hurst, J. L. 2012.
- Pheromonal induction of spatial learning in mice. *Science*, **338**, 1462-1465. doi:
- 657 10.1126/science.1225638.
- **Rosenson, L. M. & Asheroff, A. K.** 1975. Maternal aggression in CD-1 mice:
- Influence of the hormonal condition of the intruder. *Behavioral Biology*, **15**, 219-224.
- 660 doi: 10.1016/S0091-6773(75)91603-X.
- 661 **Segovia, S. & Guillamon, A.** 1993. Sexual dimorphism in the vomeronasal pathway
- and sex differences in reproductive behaviors. *Brain Research Reviews*, **18**, 51-74.
- Wells, D.J., Playle, L.C., Enser, W.E., Flecknell, P.A., Gardiner, M.A., Holland, J.,
- Howard, B.R., Hubrecht, R., Humphreys, K.R., Jackson, I.J., Lane, N.,
- Maconochie, M., Mason, G., Morton, D.B., Raymond, R., Robinson, V., Smith,
- **J.A., Watt, N.** 2006. Assessing the welfare of genetically altered mice. *Laboratory*
- 667 Animals 40:111-114.
- Wyatt, T. D. 2010. Pheromones and signature mixtures: defining species-wide signals
- and variable cues for identity in both invertebrates and vertebrates. *Journal of*
- 670 *Comparative Physiology A*, **196**, 685-700. doi: 10.1007/s00359-010-0564-y.
- 671 **Wyatt, T. D.** 2009. Fifty years of pheromones. *Nature*, **457**, 262-263. doi:
- 672 10.1038/457262a.

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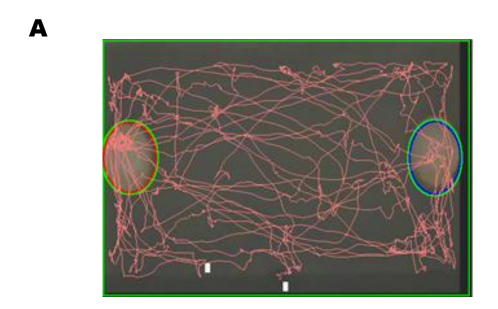
- 673 Wysocki, C. J., Wellington, J. L. & Beauchamp, G. K. 1980. Access of urinary
- nonvolatiles to the mammalian vomeronasal organ. *Science*, **207**, 781-783.

### FIGURE LEGENDS

Figure 1. **A**: Example videotrack of the exploratory behaviour of one animal in an odour preference test (citralva versus isoamyl acetate). The white pieces of scented paper are visible inside the areas of measure. **B**: Time (mean  $\pm$  SE) spent by female mice in the areas where the odorant stimuli were presented, in the experimental (left panel, darcin group) and control (right panel) groups. Grey bars in the control test represent clean pieces of paper. Orange bars represent time spent in the citralva area. Green bars represent time spent in the isoamyl acetate area. In test 2 in the experimental group, the isoamyl acetate-scented paper was also impregnated with r-darcin. Females showed a significant innate preference to remain near to the r-darcin sample (p = 0.032) and a learned preference to stay next to the isoamyl acetate-scented paper in the post training preference test (p = 0.031). In the absence of darcin, no preference appeared for any odorant.

Figure 2. **A**: Time (mean  $\pm$  SE) spent by female mice in areas where urine stimuli were presented. The left side corresponds to females that were presented with urine of healthy males versus females. Grey bars represent time spent near pieces of paper with a sample of PBS (control). The green bar represents time spent near urine of healthy males, and purple bar the time spent near female urine. The right side corresponds to females that were presented with urine of parasitized males versus females. Grey bars represent the control test. Females showed no preference for the urine of infected males (dark red bar) versus female urine (purple bar). **B**: SDS-PAGE of urinary protein for healthy male mice (n = 2), male mice parasitized with *Aspiculuris tetraptera* (n = 2), castrated male mice (n = 1) and females (n = 2). The 20 kDa band present in all cases corresponds to the molecular weight to the major urinary proteins. The small band with

- higher mobility (around 17 kDa), present in healthy and infected males, corresponds to
- the expected position of darcin.



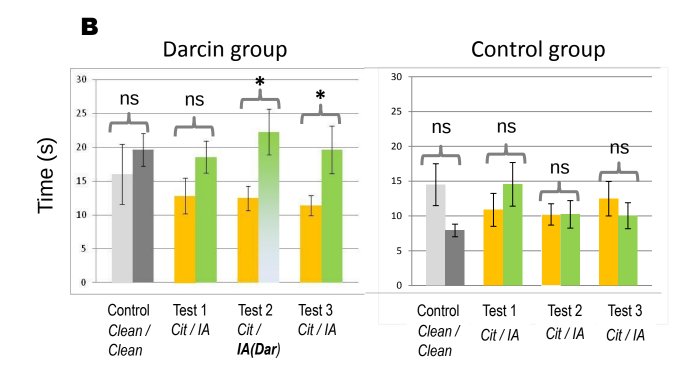


Figure 2

