1	A new test of value-modulated attentional bias in mice
2	
3	Iasmina Hornoiu, John Gigg & Deborah Talmi
4	
5	^a Division of Neuroscience and Experimental Psychology, University of Manchester
6	
7	
8	
9	
10	
11	
12	Address for correspondence
13	Deborah Talmi, Division of Neuroscience and Experimental Psychology, School of Biological
14	Sciences, University of Manchester, Manchester, UK, M139PL.
15	Telephone: 0161 275 1968
16	Email: <u>Deborah.Talmi@manchester.ac.uk</u>

Abstract

The allocation of attention can be modulated by the emotional value of the stimulus. In order to understand the biasing influence of emotion on attention allocation further, we require an animal test of value-modulated attention capture evoked by ethologically valid stimuli. In mice, female odour triggers arousal and elicits emotional responses in males. Here, we determined the extent to which objects infused with female odour captured the attention of male mice. Seven experiments were conducted, using a modified version of the spontaneous Novel Object Recognition task. Attention was operationalised using differential exploration time of identical objects that were infused with female mouse dour (O+), infused with almond odour (Oa), or not infused with any odour (O-); and non-infused novel objects (X-). We found that when single objects were presented, as well as when two objects were presented simultaneously and thus competed with each other for attention, O+ captured attention preferentially compared to O-. This was the case both when O+ were placed in a novel location and when they were placed in a familiar location. When compared with Oa at novel location, O+ at familiar location attracted more attention. Compared to X-, O+ captured more attention only when they were placed in a novel location, but attention to O+ and X- was equivalent when they were placed in a familiar location. These results demonstrate that in mice, female odour can in some circumstances capture more attention than non-ethologically relevant olfactory stimuli and object novelty. The findings of this study pave the way to using motivationally-significant odours to modulate the cognitive processes that give rise to novel object recognition. Keywords: attention, emotional arousal, memory, Novel Object Recognition task, olfactory stimuli

55

56

Introduction

57 Emotional arousal influences cogitation extensively, from early perception, to attention (Golomb, 58 Nguyen-Phuc, Mazer, McCarthy, & Chun, 2010; Pourtois, Schettino, & Vuilleumier, 2013), to higher 59 order cognitive functions (Pessoa, 2009), including memory (Levine & Edelstein, 2009; Talmi, 2013). 60 A central objective of human neuropsychological and neuroimaging research is to trace the 61 neurobiological underpinnings of the link between emotional arousal and attention (Hartikainen, 62 Ogawa, Soltani, & Knight, 2007; Kadohisa, 2013; Lee, Sakaki, Cheng, Velasco, & Mather, 2014; 63 Talmi & McGarry, 2012; Vermeulen, Godefroid, & Mermillod, 2009). In line with this objective, the 64 present study was designed to investigate the impact of an emotionally charged stimulus on attention 65 allocation in mice. Establishing the means to measure such a link would provide a valuable animal 66 task with which to further understand the neurobiology of emotional memory and attention.

67

68 Attention is crucial for the effective processing of perceptual information presented by the 69 environment at any given time (Chun, Golomb, & Turk-Browne, 2011). Behaviourally-relevant 70 stimuli are thought to be selected against others and prioritised via two routes: top-down and bottom-71 up (Corbetta, Patel, & Shulman, 2008; Miller & Cohen, 2001; Pratt & Hommel, 2003). Endogenous 72 attentional control, also referred to as *goal-directed* or *top-down*, is a voluntary process that operates 73 at the level of memory and decision making to modulate information according to the agent's 74 intentions. In contrast, the exogenous, involuntary attentional capture by certain stimuli, known as 75 bottom-up or stimulus-driven attentional process, depends on the physical characteristics of stimuli 76 and is outside the agent's control. A recently proposed model, called Multiple Attention Gain Control 77 (MAGiC), suggests a third route to attentional selection, through emotion. Emotion is thought to bias 78 perception via distinct gain control mechanisms originating in the amygdala. Neuropsychological, 79 neuroimaging and behavioural studies have provided evidence for the modulatory effects of emotion 80 on sensory processing, whereby emotionally charged stimuli acquire increased representation and 81 access to awareness compared to neutral stimuli (for review, see Pourtois et al., 2013). The majority 82 of studies that informed the MAGiC model utilised threat-related stimuli, due to their obvious 83 behavioural salience (Compton, 2003; Vuilleumier, 2005), as well as their role in various pathological 84 conditions in humans, such as anxiety and phobias (Öhman & Mineka, 2001). However, in order to 85 achieve a more complex understanding of the influence of emotion on the selection of sensory 86 information, more research is needed on the effects of other emotional stimuli on attention, including 87 those with positive value.

88

Stimuli that predict a valuable outcome – whether positive or negative - capture attention
automatically, even when they are task-irrelevant (Anderson, Laurent, & Yantis, 2011; Wang, Duan,
Theeuwes, & Zhou, 2014; Wentura, Müller, & Rothermund, 2014). This value-derived influence on

92 attention is neither top-down, nor bottom-up, since attention in this case is neither voluntarily directed 93 by contextually relevant goals, nor driven by the sensory significance of the stimuli, respectively. 94 Rather, attention is directed to stimuli that have acquired the potential to predict valuable outcomes 95 via associative learning. This mechanism is referred to as value-modulated attentional capture 96 (VMAC) (Le Pelley, Mitchell, Beesley, George, & Wills, 2016). While prioritising unexpected high-97 value stimuli might be biologically advantageous in certain situations, in others it could interfere with 98 goal-directed behaviours. Further research is needed to offer a more in-depth understanding of the 99 computational control of VMAC (Talmi, Slapkova, & Wieser, 2018).

100

101 The neural basis of top-down and bottom-up attention has been studied in detail in humans and in animal models. In humans, the majority of studies have been conducted on visual attention. 102 103 Endogenous (top-down) signals arising in higher-order prefrontal, parietal and limbic cortices interact 104 with exogenous (bottom-up) signals driven by visual cortical pathways to bias attention in favour of 105 attended targets, while suppressing representations of unattended information (Desimone & Duncan, 106 1995; Gitelman, 2003; Katsuki & Constantinidis, 2014). Across the literature, goal-directed 107 attentional processing has been generally attributed to dorsal fronto-parietal brain areas, while 108 stimulus-driven attentional control is believed to be mediated by ventral temporal-parietal networks 109 (Anderson, Laurent, & Yantis, 2014; Corbetta & Shulman, 2002; Yantis et al., 2002). Animal studies 110 of sustained attention, particularly in task-performing rats, have revealed the role of the cholinergic 111 system in top-down, as well as bottom-up attentional control. The basal forebrain corticopetal 112 cholinergic projections, which are activated via direct glutamatergic inputs from the prefrontal cortex 113 to the basal forebrain, terminate in all cortical areas and are thought to mediate top-down processes in 114 tasks involving sustained attention (McGaughy & Sarter, 1998; Sarter, Givens, & Bruno, 2001; Zaborszky, Gaykema, Swanson, & Cullinan, 1997). Lesions of the basal forebrain, affecting inputs of 115 116 acetylcholine particularly in fronto-dorsal cortical areas of rat models of attention, result in prolonged 117 impairments of sustained attention (Sarter et al., 2001). Bottom-up attentional processes have been shown to largely depend on noradrenergic projections, which originate in the locus coeruleus and 118 119 terminate in the thalamus and the basal forebrain. Noradrenergic activation of basal forebrain corticopetal projections is involved in the processing of threat-related or anxiety-inducing stimuli in a 120 121 bottom-up fashion via the recruitment of telencephalic systems (Aston-Jones, Rajkowski, Kubiak, 122 Valentino, & Shipley, 1996; Sarter et al., 2001). So far, these animal studies have demonstrated that 123 the basal forebrain cortical cholinergic system represents a core component of the neuronal circuitry 124 involved in attentional processing, thus contributing to the understanding of the role of the fronto-125 parietal cortical regions from human neuropsychological and imaging research. Integrating evidence 126 from human and animal works is necessary for the development of a reliable model of the neural 127 mechanisms of attention.

129 The neural basis of VMAC has been less well understood in animal models. In humans, Anderson et 130 al. (2014) argue that the neural mechanisms underlying VMAC are mediated by the tail of the caudate 131 nucleus and the extrastriate visual cortex. Using functional magnetic resonance imaging (fMRI), these 132 researchers found that task-irrelevant reward-predictive stimuli acting as distractors acquired stronger 133 representation in the caudate tail and triggered greater activity in the extrastriate visual cortex versus 134 other non-target stimuli. So far, animal models of attention using stimuli with emotional value have 135 been derived from the traditional theories of associative learning and conditioning. In one model of 136 associative learning, proposed by Mackintosh (1975), animals pay more attention to cues that are 137 reliable predictors of a consequence (high predictiveness) than to non-predictive cues. Selective 138 attentional bias towards good predictors allows animals to focus on relevant cues, while ignoring 139 distractors, thereby achieving optimal performance. In contrast, the model by Pearce and Hall (1980) 140 states that cues with uncertain consequences capture the most attention. The idea behind this model is 141 that because unreliable cues are surprising, they will be allocated more attentional resources that leads 142 to rapid learning about their significance. While there is abundant evidence in favour of both 143 predictability (Duffaud, Killcross, & George, 2007; George & Pearce, 1999) and uncertainty (Kaye & 144 Pearce, 1984; Wilson, Boumphrey, & Pearce, 1992) models, several hybrid models have emerged in 145 an attempt to reconcile these principles (for reviews, see Esber & Haselgrove, 2011; Le Pelley et al., 146 2016). To date, very little, if any, data exist on the influence of biologically relevant stimuli on 147 VMAC in rodents. The present study is, to our knowledge, the first research using female odours to 148 examine VMAC in a rodent model of attention.

149

150 Unlike humans, who are predominantly influenced by visual stimuli (Shapiro, Egerman, & Klein, 151 1984), most mammals, including rodents, rely mainly on odours to obtain information about their 152 environment (Brennan & Kendrick, 2006; Johnston, 2003). Odours from conspecifics of opposite sex 153 are thought to carry positive reward values, because they elicit an approach response to promote 154 reproductive behaviours, as opposed to withdrawal or avoidance responses, which supress 155 reproductive behaviours (Beny & Kimchi, 2014). The motivational significance of odours bearing 156 reproductive value does not seem to depend on the animal's prior sexual experience; these stimuli 157 can, therefore, be considered primary reinforcers. This is evident in the laboratory, where sexually 158 inexperienced male rodents display typical sexual behaviours when exposed to female odours (Beny 159 & Kimchi, 2014). Given the socio-biological relevance of odours and their known motivational 160 effects on animals, it is reasonable to assume that odours could be used as stimuli to investigate 161 VMAC in an animal model.

162

163 Seven experiments were designed to quantify the degree of attention allocation in male mice to 164 objects infused with female odour (O+). We contrasted the O+ objects with odour-neutral objects (O-165), novel objects (X-) and objects infused with almond extract (O^a), a mildly attractive odour for

166 rodents (Huckins, Logan, & Sanchez-Andrade, 2013). These objects were used in a modified version 167 of the spontaneous Novel Object Recognition (NOR) task. NOR is one of the most widely used paradigms in studies of memory functions in rodents. The NOR task can be configured to measure 168 169 attention (Silvers, Harrod, Mactutus, & Booze, 2007) and has been used in studies focusing on 170 pathological conditions or drug abuse, which result in attentional deficits (Alkam et al., 2011; Piper, 171 Fraiman, & Meyer, 2005). Originally developed by Berlyne (1950) and subsequently adapted for use 172 in mice by Murai et al. (2007), this behavioural task relies on the drive of rodents to explore novelty 173 in the absence of any training or external reinforcers. The original NOR task comprises three phases: 174 habituation, familiarisation and test. In the first two phases, the animal is exposed to an open-field 175 arena (habituation) and then to the same arena in the presence of two identical objects 176 (familiarisation). The test phase is similar to the familiarisation phase, with the exception that one of 177 the identical objects is replaced with a novel object. Healthy adult rodents recognise the familiar 178 object at test and pay more attention to the novel (more arousing) object, indicated by a longer 179 exploration time of the novel versus familiar object (Ennaceur, 2010; Gaskin et al., 2010; Mumby, 180 Glenn, Nesbitt, & Kyriazis, 2002). In the modified NOR paradigm used in this study, instead of novel 181 and familiar objects, the attention allocation of male mice was investigated for O+ versus O-, O^a or X-182 , which were placed in either novel or familiar locations in a Y-maze.

183

184 The first two experiments assessed the attentional capture of mice by an O+ compared with an O, 185 when both objects were novel (Experiment 1) or familiar (Experiment 2). We hypothesised that 186 because the motivational value of O+ was higher than of O-, the former would attract more attention 187 in both experiments. We then asked whether the same would be true if O+ and O- were placed at 188 different locations in the arena, so that O+ would be found at either a novel (Experiment 3) or a 189 familiar (Experiment 4) location. In these experiments, novel location was used due to its previously 190 demonstrated influence on attention (Ennaceur, Neave, & Aggleton, 1997) and, thus, it was 191 interesting to study how well novel location competed with female odour for attention allocation. We 192 hypothesised that when the location of O+ was novel, O+ would capture more attention than O-, 193 because the former benefited from the advantage of two salient motivational features, namely, motivational value and novel location. However, in Experiment 4 it was less clear whether O+ would 194 still 'win' over O-, due to the fact that the novel location of the latter was now competing with the 195 196 odour of the former for attention. In the next two experiments, we used another well-known 197 motivational feature, novel object identity, and assessed how strong the attention capture by odour 198 was when competing with a novel object (X-). Novel objects are expected to recruit more attention 199 than familiar objects in the traditional NOR task and, therefore, we were interested to test how 200 attention to O+ would be influenced by the presence of X-. Experiment 5 tested the attention captured by O+ at a novel location versus an X- at a familiar location. Here, O+ had the advantage of odour 201 202 and location novelty, while X- had the advantage of novel identity. In Experiment 6, X- was placed at

a novel location and O+ at a familiar location. Since this time X- had the advantage of both novel
 identity and novel location, the question was whether it would attract more attention than O+.

205

206 There is preliminary evidence that social odour captures more attention in mice compared to non-207 social odours. Rattazzi, Cariboni, Poojara, Shoenfeld, & D'Acquisto (2015) compared female mouse 208 odour and other social odours to non-social odours in male mice of a similar strain to the ones used 209 here, and which were obtained from the same provider. They found that control mice, as well as 210 immune-cell deficient recombination-activating gene (RAG-1) knockout mice, showed increased 211 sniffing time for the former. Therefore, in Experiment 7, we wanted to compare the degree to which 212 objects infused with female odour attracted more attention than objects infused with another odour 213 with a lower level of motivational significance.

214

215 In their study on the effects of odour preferences on rats' discrimination learning, Devore, Lee, & 216 Linster (2013) classified 53 monomolecular odorants as high, neutral and low, in terms of 217 spontaneous odour preferences. By contrast, the odour of flowers, nuts and fruit is a mixture of 218 compounds. For the purposes of our study, instead of single chemical compounds, we decided to use 219 the odour from a compound mixture (such as a flower, nut or fruit), because such odours are prevalent 220 in behavioural studies of mice (Arbuckle, Smith, Gomez, & Lugo, 2015; Rattazzi, Cariboni, Poojara, 221 Shenfeld, & D'Acquisto, 2015; Yang & Crawley, 2010). We selected almond odour, one of those 222 used by Rattazzi et al. (2015). According to (Huckins, Logan, & Sanchez-Andrade, 2013), unlike 223 conspecific urine smell, which is a 'social odour' and, therefore, high in motivational significance, the 224 odour of almond extract is mildly motivationally significant, since it is a natural food odour but 225 distinct from the food laboratory mice are used to. This type of odour can be employed as a neutral 226 non-social odour alongside social odours in rodent experiments investigating odour-mediated 227 behaviour and odour identification ability (for instance, when testing deficits in identifying pleasant or 228 neutral odours in rodent models of psychiatric diseases, Huckins, Logan, & Sanchez-Andrade, 2013). 229 We predicted that despite the mild motivational significance of almond odour, animals in Experiment 230 7 will allocate more attention to female mouse odour.

231

In Experiment 7b attention allocation to O+ at a familiar location was compared to attention allocation to O^a at a novel location. Here, O+ had the advantage of motivational significance, while O^a benefited from two factors, namely location novelty and a mildly attractive odour. This experiment compared the attentional draw of two different odours, in contrast with the previous two experiments, which compared an olfactory stimulus with visual stimuli. Similar to Experiments 5 and 6, in this last experiment it was interesting to observe which object feature combinations captured more attention and whether O+ can still elicit more interest.

240	Unlike previous research which used social and non-social odours to look at the animal's capacity to
241	distinguish between the two and to investigate habituation/dishabituation (Arbuckle, Smith, Gomez,
242	& Lugo, 2015; Yang & Crawley, 2010), or to test altered sense of smell in certain conditions
243	(Rattazzi, Cariboni, Poojara, Shenfeld, & D'Acquisto, 2015), our study focused on quantifying the
244	attentional capture by a social odour and provided a reliable protocol for future studies using NOR in
245	such behavioural assessments.
246	
247	Methods
248	
249	Animals
250	Behavioural experiments were performed using eight adult male C57BL/6J mice (Charles River, UK),
251	which were 14 weeks (3.5 months) old at the start of the experiments and weighed 30±3 g throughout
252	the project. Mice were weighed prior to each experiment and at the end of the last experiment on a
253	laboratory scale (Kern FCB, Germany). Mice were maintained by the Biological Services Facility
254	(BSF), University of Manchester, UK and housed in groups of four individuals in ventilated
255	Techniplast cages, in standard conditions (20°± 2°C temperature and 55±5% humidity) on a 12:12

Behavioural experiments were performed using eight adult male C57BL/6J mice (Charles River, UK), which were 14 weeks (3.5 months) old at the start of the experiments and weighed 30 ± 3 g throughout the project. Mice were weighed prior to each experiment and at the end of the last experiment on a laboratory scale (Kern FCB, Germany). Mice were maintained by the Biological Services Facility (BSF), University of Manchester, UK and housed in groups of four individuals in ventilated Techniplast cages, in standard conditions ($20^{\circ}\pm 2^{\circ}$ C temperature and $55\pm5\%$ humidity) on a 12:12 light/dark cycle, with *ad libitum* access to food and water. All mice were ear punched for identification. The experiments were carried out over a period of two months and took place between 9:30am-12:30pm. All procedures were conducted in conformity with the University of Manchester BSF regulations for animal husbandry and with the Home Office Animals (Scientific Procedures) Act (1986) and were licenced by the UK Home Office and University of Manchester Ethical Review Panel.

262

263 Apparatus

264 Experiments were performed in custom Y-mazes with three identical white, opaque plastic arms, 265 (length 160mm, height 280mm) diverging at a 120° angle from each other (designed by Jack Rivers-Auty and constructed by Plastic Formers Ltd, UK). Each arm became wider at the end to form a small 266 square arena (length 92mm x width 90mm). The square arenas could be differentiated by the presence 267 268 of salient visual cues. Individual mice were randomly assigned to a particular Y-maze throughout 269 habituation and testing. Mice were always released inside the middle arm (the arm closest to the 270 nearest room wall), with their backs to the right and left arms, which contained objects (see 271 Materials). This strategy, following the guidelines by Antunes and Biala (2012), ensured that external 272 pressure to explore the objects was avoided. During the interval between exposures (habituation) and 273 the inter-trial interval (ITI, experiments), mice from each cage group were placed in separate holding 274 cages (standard housing cages, Techniplast), which remained the same throughout the project. Mice

from each cage group were run simultaneously in four Y-mazes placed next to each other. Videocameras (JVC, 40x optical zoom) placed above the Y-mazes recorded animal performance.

277

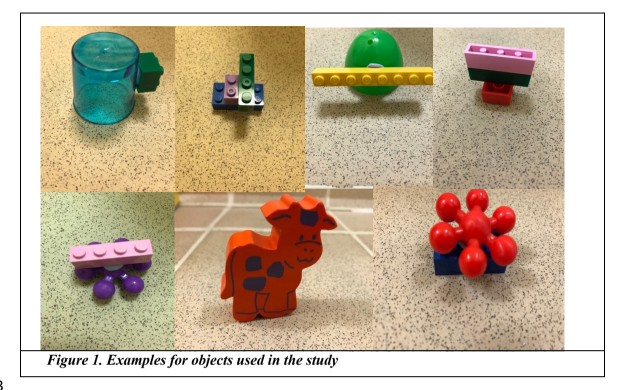
278

279 Materials

The objects used in experiments were either built from LEGO[®] pieces or various other plastic shapes 280 281 (Figure 1). Given that the egg halves were identical in shape, the difference in their colour as well as 282 their position in the maze (face-down or on one side with either convex or concave side facing the 283 animal) were used to create different object types. All objects were odour-neutral and made of plastic 284 in order to avoid material preference, minimise odour saturation and facilitate cleaning. The objects were attached to the floor inside the Y-maze with Blu-Tack^{\mathbb{R}}. Objects used in the habituation phase 285 were plastic letters and were never used in subsequent experimental trials. New object types were 286 287 used in each experiment, in order to avoid habituation to any object type.

288

289 For each experiment, soiled cage bedding from cages containing four female mice (C57BL/6J strain) 290 was used to label some objects with female odour. The objects to be labelled with female odour were 291 placed in the bedding the day prior to each trial, around midday (12:00pm). For the experiments that 292 lasted four days (Experiments 1, 3 and 4), halfway through each experiment (at the end of day 2), the 293 bedding with female odour was replaced with bedding from a cage containing a second group of 294 control females of the same strain. For the experiments that lasted only 2 days (Experiments 2, 5 and 295 6), the bedding remained the same throughout each experiment, but was changed before the start of 296 the next experiment. The reason for this was that it was observed that the mice had a tendency to habituate to the smell after day 2 and using new bedding prevented this. Objects with female odour 297 298 are here referred to as 'O+'. Copies of the same object but without odour are referred to as 'O-'. 299 Objects that were never infused with odour are termed 'X-'. Y-mazes and objects were cleaned 300 thoroughly with 70% ethanol and wiped with paper towels between trials with mice from different 301 group cages, as well as at the end of each daily session.



303

304 Procedure. The tasks used in this study were a modified version of the classical NOR task (Berlyne,
305 1950). The procedure of each experiment is depicted in Figure 2. The study began with a
306 familiarisation stage, during which the animals were handled for one minute on two consecutive days
307 so that they became accustomed to the experimenter.

308

309 Habituation. After familiarisation, the mice were habituated to the testing apparatus and objects over 310 a five-day period prior to Experiment 1. The first day of habituation consisted of a 10-minute cage 311 group habituation session to the Y-mazes, in which mice from a given cage were placed together in 312 one Y-maze, in the absence of objects. On the following day, the mice were exposed individually to 313 the Y-maze for 10 minutes, again without objects. On the third day, the same steps from day 2 were 314 repeated, but this time an O was present in either the left or right choice arm. On the fourth day, 315 following the same steps as on day 3, each animal was exposed to another O-, but in the opposite arm 316 to the one on day 3. The last day of habituation consisted of two exposures, this time with an O+ in 317 either the left or right arm of the maze. The duration of exposure was 10 minutes and the interval 318 between exposures was 5 minutes. During the 5-minute interval, the mice were placed in their holding 319 cages and the O+s presented in the first exposure were replaced with different O+s for the second 320 exposure. In addition, mice were habituated to O^a objects in either left or right arm.

321

322 Experimental stage. Following habituation, the mice were tested in six different experiments.
323 Experiments 1, 3 and 4 were conducted on four consecutive days. Experiments 2, 5, 6 and 7b were
324 conducted on two consecutive days. There were two experimental trials each day. The interval

325 between two experimental trials (ITI) was 5 minutes. During the ITI, the animals were placed in 326 holding cages and the objects were replaced for the next trial. The particular object used in a given 327 trial (O+, O-, X- or O^a), the order in which objects were presented and their locations in the maze (left 328 or right choice arm) were randomised for each animal in each experiment. This randomisation 329 ensured that each animal was exposed to a given object only in one experimental trial across the entire 330 study. Each animal experienced the same number of O+, O-, O^a and X objects in each experiment. 331 Apart from Experiment 1, each animal was exposed only to one O+ on each day and whether this was 332 in the first or second trial of the day was counterbalanced. All of the experiments were performed 333 consecutively by the same animals. Chronologically, the experiments were carried out in the 334 following order: Experiment 1, Experiment 3, Experiment 4, Experiment 2, Experiment 5, 335 Experiment 6, Experiments 7a and 7b. Thus, the mice were 14 weeks (3.5 months) old in Experiment 336 1, 16 weeks (4 months) old in Experiment 3, 19 weeks (almost 5 months) old in Experiment 4 and 29-337 30 weeks (around 7.5 months) old in Experiments 2, 5, 6 and 7a and 7b.

338

Experiment 1. This experiment consisted of eight trials in total per animal. Trials included one
exposure to a single object, either O+ or O-. The duration of each trial was one minute (see row 1 of
Fig. 2).

342

Experiment 2. This experiment was composed of four trials per animal. Each trial included an initial
exposure to O- for one minute (the 'study phase'), either in the left or the right arm of the maze,
followed by an exposure either to O- or O+ for 3 minutes (the 'test phase'; see row 2 of Fig. 2).

346

Experiment 3. This experiment included eight trials in total per animal. Each trial included a study phase, where animals were exposed to O- for one minute, either in the left or the right arm of the maze. The location of O- in the study phase is referred to as the 'familiar' location. The other location - the arm that was empty during the study phase – is referred to as the 'novel' location. During the test phase animals were exposed to both O- and O+ for 3 minutes. The O+ was always placed in the 'novel' location (see row 3 of Fig. 2).

353

354 Experiment 4. This experiment was identical to experiment 2, except that in the test phase O+ was in355 a familiar location (see row 4 of Fig. 2).

356

Experiment 5. This experiment was composed of four trials in total per animal. Each trial included a
study phase, where animals were exposed to O- for one minute, either in the left or the right arm of
the maze. The location of O- in the study phase is referred to as the 'familiar' location. The other
location – the arm that was empty during the study phase – is referred to as the 'novel' location.
During the test phase animals were exposed to both O+ and X- (an entirely novel object) for 3

362 minutes. The O+ was always placed in the 'novel' location (see row 5 of Fig. 2).

363

364 Experiment 6. This experiment was identical to experiment 5, except that O+ in the test phase always365 occupied a familiar location (see row 6 of Fig. 2).

366

Experiment 7a. In order to ensure that the mice were to some extent attracted to almond odour, we tested the preference of mice for almond odour in a four-trial-per-animal experiment, where mice were simultaneously exposed to two filter papers, one without any odour and the other infused with almond smell, at randomly determined locations in the Y-maze (left or right arm). To label the almond-odour paper with almond smell, we used a cotton-tipped applicator dipped in pure almond oil (100% concentration, by Atlantic Aromatics, Bray, Co.Wicklow, Ireland) and then scrubbed it on a small piece of filter paper.

374

375 **Experiment 7b.** In this experiment, O+ in the test phase was always placed at a familiar location and 376 the O^a occupied a novel location (see row 7 of Fig. 2). The number of trials and the way the 377 experiment was conducted was identical to Experiments 5 and 6. For labelling the objects with 378 almond odour, we scrubbed O objects with cotton-tipped applicators dipped in pure almond oil (100% 379 concentration, by Atlantic Aromatics, Bray, Co.Wicklow, Ireland), thus obtaining O^as; in addition, in 380 order to remove the oiliness from these objects which could have led to a possible bias in attention 381 capture (arising from a difference in texture between O+ and O-), the latter were also gently wiped 382 with paper towel before being placed in the Y-mazes.

- 383
- 384
- 385
- 386
- 387
- 388
- 389
- 390
- 391
- 392
- 393
- 394
- 395
- 396
- 397
- 398

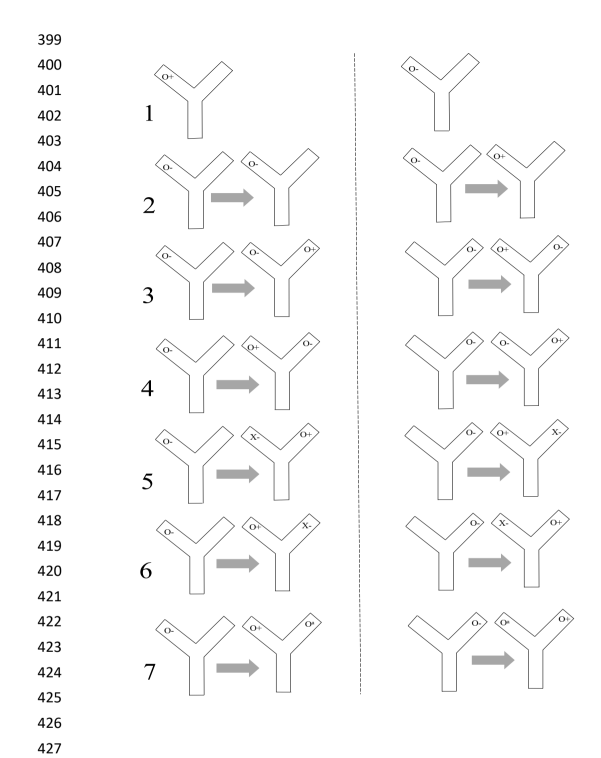


Figure 2. Experimental design.

Trial types in experiments 1-7 are illustrated (one experiment in each row). O+: female-mouseodour-infused object. O-: odour-neutral object. X-: novel object, never infused with odour. O^a: almondodour-infused object. Different copies of objects were used in two-phase experiments (Experiments 3-7).

436 Data analysis. Exploratory behaviour was recorded with video cameras and subsequently, object 437 exploration time for each object was scored using the Novel Object Timer software (Jack Rivers-438 Auty; Novel Object Recognition Task Timer, 2015). The animal was considered to be exploring an 439 object when its nose was within 2cm of the object and directed at the object. Sitting next to the object 440 or climbing on top of the object was not regarded as exploratory behaviour. Each trial was scored 441 twice for accuracy and the average of the two scorings taken as the object exploration time per trial. 442 The mean exploration time of O+, O-, O^a and X- was obtained by averaging all trials with these 443 objects for each animal.

444

The displacement index (D2), used to assess object preference from Experiments 3 onwards, was calculated using the formula: $D2 = (T_{O^+} - T_{O^-}) / (T_{O^+} + T_{O^-})$, where T_{O^+} is the mean exploration time of O+ objects and T_{O^-} is the mean exploration time of O objects. The values for this index are bound within a range of -1 and 1; positive D2 values indicated preference for O+, while negative values signified preference for O- and a value of 0 signified no preference (Burke, Wallace, Nematollahi, Uprety, & Barnes, 2010; Oliveira, Hawk, Abel, & Havekes, 2010).

451

The differences in total exploration time of O^+ , O^- , O^a and X- were analysed with paired t-tests (twotailed). D2 values were analysed with one-sample t-tests. In Experiments 3-4, we also investigated the effect of experiment day, using a repeated-measures 2 (Object type: O^+ , O^-) x 4 (day: 1-4) ANOVA.

- 455 Statistical tests were performed in GraphPad Prism 8.
- 456

457

458

Results

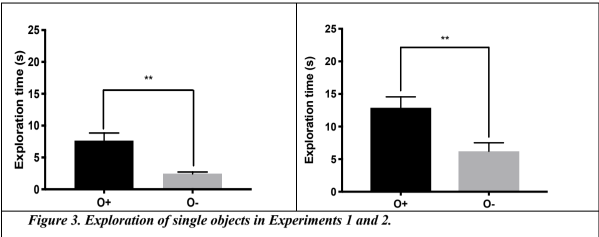
459 Experiment 1: Novel objects O+ vs. O-

460 The total time animals spent exploring O+ was significantly higher than the total exploration time for 461 O- (t(5)=4.11, p=0.0092) as illustrated in Figure 3 (left panel). This finding demonstrates that when 462 both O+ and O- are novel, the former attracts more attention. Two animals were excluded from the 463 analysis of Experiment 1 due to a counterbalancing error.

464

465 Experiment 2: Familiar objects O+ vs. O-

466 The total object exploration time in Experiment 2 (Fig. 3 right panel) was significantly higher for O+ 467 than O- (t(7)=3.67, p=0.008). This finding demonstrates that when both O+ and O- are familiar, the 468 former attracts more attention.



Left: Results of Experiment 1: Novel O+ vs. Novel O-. *Right:* Results of Experiment 2: Familiar O+ vs. Familiar O-. Bar graphs represent object exploration times (in seconds) averaged across animals. Data are presented as means \pm SEM. Statistical significance is expressed with ** ($p \le 0.01$).

469

470 Experiment 3: Competition between O+ in a novel location and O- in a familiar location.

471 Figure 4 shows that the total exploration time for O+ was significantly higher than that for O-(t(7)=6.99, p=0.0002). Additionally, animals demonstrated a significant preference for O+ over O-, 472 473 as indicated by D2 value, which was significantly higher than zero (t(7)=9.33, p<0.0001). Object 474 exploration time was analysed across experimental days for any effect of day or interaction between 475 object type and day. A repeated-measures two-way-ANOVA test found that there was a significant 476 effect of object type (F(1,7)=47.84, p<0.001). While numerically this difference diminished across 477 days, neither the effect of day (F(3,21)=2.93, p=0.06), nor the interaction (F(3,21)<1, p=0.53) were 478 significant.

479

480 Experiment 4: Competition between O+ in a familiar location and O- in a novel location.

The total exploration time of O+ was significantly higher than that of O- (t(7)=3.48, p=0.01; Figure 482 4). The D2 value (Figure 4), indicating the preference for O+ over O-, was significantly higher than 483 zero (t(7)=3.67, p=0.008). Object exploration time was analysed across experimental days for any day 484 effect or interaction between object type and day. Replicating the above results, there was a 485 significant effect of object type (F(1,7)=12.02, p=0.01), as determined by a two-way-ANOVA test. As 486 in the previous experiment, here the effect of day (F(3,21)=2.24, p=0.11) and the interaction 487 (F(3,21)=1.13, p=0.36) were not significant.

488

At the end of the third experiment, it was interesting to determine if the mice had higher preference
for O+ placed in a novel location than for O+ in a familiar location. The overall D2 values from
experiments 3 and 4 were statistically compared with a Student's paired t-test. As illustrated in Figure

492 4, the D2 value for O+ in a novel location was significantly greater than that for O+ in a familiar
location (t(7)=2.32, p=0.027), indicating that location novelty increased the motivational value of O+.
However, since the D2 values were from two separate experiments, it is important to take into account
possible confounds arising from a counter effect of habituation. Since Experiment 4 followed
Experiment 3, animals might have been less motivated to explore the objects in the former than in the
latter.

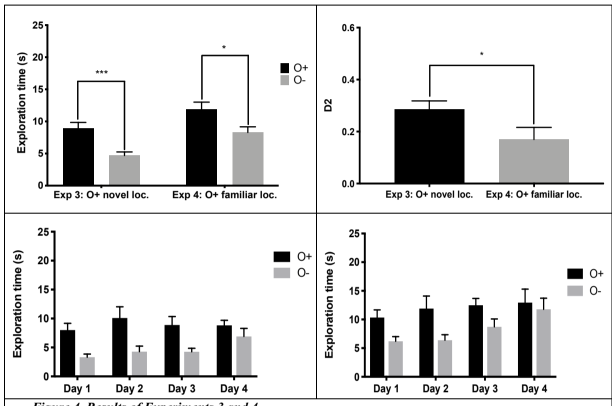


Figure 4. Results of Experiments 3 and 4.

Top left: Object exploration time in Experiments 3 and 4. Means of object exploration time (seconds) were calculated by averaging across all animals.

Top right: *Effect of location familiarity on odour preference. Mean D2 (discrimination index) value was obtained as the average of the D2 values from all animals.*

Bottom: Daily object exploration in Experiment 3 (Left) and 4 (Right). Means of object exploration time (seconds) were obtained by averaging across all animals for each experimental day. The bar graphs represent means \pm SEM. Statistical significance is expressed as ns (p>0.05), * (p≤0.05) or *** (p≤0.001).

498

499 Experiment 5: Competition between O+ in a novel location and X- in a familiar location.

- 500 As shown in Figure 5, the total object exploration time in Experiment 5 was significantly higher for
- 501 O+ than for X (t(7)=5.63, p=0.0008). A one sample t-test showed that the D2 value was significantly
- 502 higher than zero, indicating a preference for O+ over X (t(7)=4.92, p=0.0017).
- 503

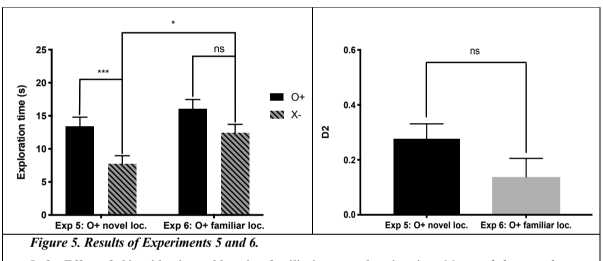
504 Experiment 6: Competition between O+ in a familiar location and X- in a novel location.

505 The total object exploration time in Experiment 6 was higher for O+ than for X-, however, the

506 difference was not statistically significant (t(7)=1.8, p=0.1158, Figure 5). The D2 value in this 507 experiment was also not significantly higher than zero, suggesting no significant preference for O+ 508 over X- (t(7)=2.023, p=0.0828).

509

510 Additionally, the results from Experiment 5 and Experiment 6 were compared in order to determine 511 the effect of location familiarity and object type (novel identity or odour-infused) on exploration time 512 and object preference. The same caveats hold for this comparison across experiments as for the one 513 across Experiments 3 and 4. A repeated-measures two-way-ANOVA test found that there was a 514 significant effect of location (F(1,7)=19.23, p=0.003) with a preference for the novel object, as well as 515 of object type (F(1,7)=11.68, p=0.011), but no significant interaction (F(1,7)=1.59, p=0.246). It is 516 difficult to interpret these results conclusively as indicative of additive or sub-additive effects, given 517 the lack of counterbalance between experiments and potential effects of habituation.



Left: Effect of object identity and location familiarity on exploration time. Means of object exploration time (seconds) were calculated by averaging across all animals. Bar graphs represent object exploration times (in seconds) averaged across animals.

Right: Effect of object identity and location familiarity on object preference. Mean D2 (discrimination index) value was obtained as the average of the D2 values from all animals. Data are presented as means \pm SEM. Statistical significance is expressed as ns (p > 0.05),* ($p \le 0.05$) or *** ($p \le 0.001$).

518

519 In Experiments 5 and 6, it was also interesting to observe the choice the animal made when it had to 520 decide which object to explore first (O+ or X-); this is referred to as *first object choice*. A binomial 521 test revealed that there was not enough evidence to reject the null hypothesis, according to which 522 animals were equally likely to explore either object (p=0.1402 in Experiment 5, p=0.2153 in 523 Experiment 6).

524

525 Experiment 7a was necessary to investigate if mice are able to detect the smell of almond and,526 moreover, if they find this smell attractive. Figure 6 shows that the total exploration time of filter

- 527 papers with almond odour was significantly greater than the exploration time of an odour-free filter 528 paper (t(7)=2.9, p=0.0229). The D2 value, which was found to be significantly higher than zero 529 (t(7)=3.5, p=0.0101), confirms the preference of mice for almond odour compared to no odour.
- 530

531 Experiment 7b: Competition between O+ in a familiar location and O^a in a novel location.

- 532 The overall exploration time for O+ was significantly higher than that for O^{a} (t(7)=7.5, p=0.0001) and
- the D2 value was significantly larger than zero (t(7)=11.5, p<0.0001, Figure), indicating preference
- 534 for O^+ over O^a .

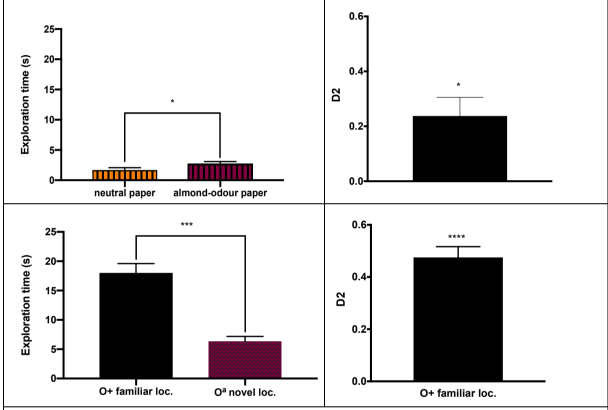


Figure 6. Results of Experiment 7.

Top left: Effect of odour on exploration time in Experiment 7a. Means of object exploration time (seconds) were calculated by averaging across all animals. Bar graphs represent exploration times (in seconds) averaged across animals.

Top right: Odour preference in Experiment 7a. Mean D2 (discrimination index) value was obtained as the average of the D2 values from all animals. Data are presented as means \pm SEM. Statistical significance is expressed as * (p<0.05).

Bottom left: Effect of odour on exploration time in Experiment 7b. Means of object exploration time (seconds) were calculated by averaging across all animals. Bar graphs represent object exploration times (in seconds) averaged across animals.

Bottom right: Effect of odour on object preference in Experiment 7b. Mean D2 (discrimination index) value was obtained as the average of the D2 values from all animals. Data are presented as means \pm SEM. Statistical significance is expressed as *** ($p \le 0.001$) or **** ($p \le 0.0001$).

535 536

Discussion

537 Evidence for VMAC from sensory domains outside vision is sparse, despite abundant research on 538 different sensory modalities involved in bottom-up and top-down attention processes (Spence, 2010). 539 The reason for this is that the initial identification of VMAC was for visual cues and since then, the 540 scientific focus has been on analysing how learnt value affects visual attention (Chelazzi et al., 2014; 541 Failing & Theeuwes, 2014; Oi, Zeng, Ding, & Li, 2013). Extending the principles of VMAC to other 542 sensory domains is essential for a more complex understanding of this attentional process and can 543 help integrate current knowledge of the modulatory effects of learnt reward on sensory processing 544 (Pantoja et al., 2007). In the study we report here, we investigated the attentional capture by an 545 olfactory stimulus and the possibility of any cross-modal interference with visual stimulation by an 546 object at a novel location or a novel object in the arena.

547

548 Previous studies have shown that female odour represents a positive arousing stimulus for laboratory 549 male mice (Beny & Kimchi, 2014; Connor, 1972; Mackintosh, 1970). According to Beny and Kimchi 550 (2014), sexually inexperienced (naïve) male mice display sexual behaviours towards female 551 conspecifics and manifest aggression towards other male mice. Connor (1972) demonstrated that 552 pheromones found in urine are responsible for these dimorphic behaviours. In his experiments, male 553 mice displayed milder aggressive behaviours towards male intruders smelling of female urine and 554 behaved aggressively towards females swabbed with male urine. These behaviours are genetically 555 determined and, therefore, do not require prior learning. The only experience these laboratory mice 556 had with female odours was during weaning, in the presence of their mothers, after which they were 557 isolated from female conspecifics.

558

In line with these observations, the results from our Experiments 1 and 2 support the conclusion that male mice pay significantly more attention to O+ than O-. In general, in these experiments, there seems to be longer exploration of familiar objects compared to novel ones, which might be explained be a slight neophobia in the mice. Taken together, these experiments confirmed our hypothesis that female odour captures more attention than an odourless object. These experiments ensured that our mice displayed behaviours as expected, based on the aforementioned literature, so we were able to proceed to the next stages of the study.

566

The following aims were to elucidate how much attention odour would capture when the location of O+ was either novel or familiar. Previous studies have already established that under normal conditions, adult rats show preference for an object at a novel location compared to an identical object at a previously experienced location (Aggleton, Albasser, Aggleton, Poirier, & Pearce, 2010; Barker & Warburton, 2011; Ennaceur et al., 1997). Therefore, it was not surprising that in Experiment 3, O+

- 572 at a novel location attracted more attention than an O- at a familiar location, since both its odour and 573 location provided O+ with more motivational significance than O-. Interestingly, in Experiment 4, 574 where O+ at a familiar location competed with O- at a novel location, O+ still attracted more 575 attention. This represents a notable finding, as it shows that female odour 'wins' over a previously 576 known arousing factor (location novelty) in the amount of attention captured. In a human study of 577 auditory attention, Anderson (2016) demonstrated that VMAC by sounds previously associated with a 578 reward interfere with a visual task and compete with the visual representation of stimuli, reflecting 579 cross-modal stimulus competition bias by VMAC. Our study shows that, at least in mice performing a 580 NOR task, a positively stimulating odour outcompetes a visual stimulus (location novelty) for 581 attention, thus extending the findings from humans to animals and also to a different sensory domain.
- 582

583 At first glance, the observation that O^+ at a novel location attracts more attention than O^+ at a 584 familiar location might seem intuitive: in the first case, O+ consists of two motivational stimuli -585 odour and location novelty, while in the second case O+ only has the odour. However, this also 586 emphasises an aspect worth taking into consideration – the fact that multiple stimuli can act in 587 combination to influence attention. Studies focusing on episodic memory have demonstrated that rats 588 and mice form integrated representations of three distinct object features: its identity, location and the 589 context in which it was experienced. Being able to associate these separate components allows the 590 animal to achieve a complex representation and record of environmental experiences, and this has 591 been termed 'episodic-like memory' in rodent models (Davis, Eacott, Easton, & Gigg, 2013; Eacott, 592 2004). In light of this work, the interpretation of how O+ at familiar location versus O+ at novel 593 location affects attention allocation could be further extended. The combination of odour and location 594 provides the animal with more detailed information about the object and, thus, it attracts more 595 attention than either odour or location alone. This suggests that odour can act not only on its own, but 596 also as a component of a stimulating context to modify the degree of attention allocation, thus 597 influencing other cognitive processes (e.g., episodic memory) in an additive manner.

598

599 Statistically, there was no day effect on the exploration time of O+ and O- in Experiments 3 and 4. 600 However, the daily means of exploration time indicate that there was a tendency for the attention 601 capture by O+ to decline across trials. This might be due to the male mice learning over the first three 602 trials that if they encounter female olfactory cues, this does not predict the availability of the female 603 mice, so the female odour starts to lose attentional allocation. This observation should be considered 604 in future behavioural studies using odours, particularly if such studies involve several trials. In our 605 study, using odour from different females after four consecutive trials was an attempt to avoid 606 habituation to odour.

608 Once we were able to determine that odour elicits more attentional capture than location novelty, the 609 next logical step was to study attention allocation to odour when competing with another powerfully 610 arousing stimulus – novel identity. The motivational value of novel identity has been the premise of 611 many studies using the NOR task, in which rodents are expected to pay more attention to the novel 612 rather than the familiar object. In our study, odour captured more attention than a novel object when 613 the location of the former was novel, while the location of the latter was familiar (Experiment 5). If 614 we assume that location is a visual stimulus, then odour in association with a visually arousing 615 stimulus captures more attention than another visual stimulus known to be salient to rodents. 616 However, in Experiment 6, when X- was placed at a novel location and O+ at a familiar location, 617 odour could no longer 'win' over novel identity plus location in the degree of attention allocation. 618 These findings suggest that, on their own, odour and object identity might have the same motivational 619 significance to mice and their combination with other arousing stimuli can shift the balance in favour 620 of one over the other.

621

622 Nevertheless, it is not yet clear how visual and olfactory stimuli interact to capture the attention of 623 mice and whether the two kinds of stimuli are indeed equally motivational. A rather surprising 624 observation in Experiment 5 was that, in a single case, a particular X- type attracted more attention 625 than its O+ counterpart; this indicates that in some circumstances, visual cues elicit stronger effects on 626 attention than olfactory cues. Le Pelley et al. (2016) argue that competition between a physically 627 salient and a less salient cue can alter attention, which can in turn affect the degree of learning. Since 628 both visual and olfactory cues convey important adaptive information to rodents, the question we 629 might ask is what determines whether a certain visual or olfactory cue attracts more attention than 630 other motivational stimuli present in the same environment. This question should motivate further 631 research into object type preference in mice, as well as more comparisons between the effects of 632 visual and olfactory cues on attention allocation.

633

634 A recent study in monkeys showed that for the visual system, novelty enhanced the motivational 635 value of stimuli associated with a negative outcome, while for the reward system, the effects of novelty dissipated for such stimuli (Foley, Jangraw, Peck, & Gottlieb, 2014). This led the researchers 636 637 to conclude that novelty acts on attentional mechanisms independent of reward to influence the 638 processing of information and learning. In light of this finding, our experiments in mice could be 639 interpreted not just in terms of competition between novelty and a reward-associated stimulus, but 640 rather as evidence that reward and novelty are dissociable. In the future, it would be interesting to 641 investigate in humans whether this distinction can be made at both psychological and neurobiological 642 levels, especially since, contrary to this theory, prior research on animals has suggested that the 643 effects of novelty can be explained in terms of reward (Horvitz, 2000; Kakade & Dayan, 2002; 644 Laurent, 2008).

645 Several studies have demonstrated that the value of a particular stimulus feature (e.g., colour) in 646 predicting reward results in more attention being allocated to the same or similar features (Kalish & 647 Kruschke, 2000; Lawrence, Sahakian, Rogers, Hodges, & Robbins, 1999; Sutherland & Mackintosh, 648 1971). In our study, male mice were likely to evaluate female odour as an endogenously highly 649 motivational stimulus. We are still uncertain as to whether naïve males consider female odour to be 650 reminiscent of their mothers and, thus, have a predictive value acquired through learning. In the 651 future, it would be useful to compare the degree of attention allocation to female odours with that to 652 odours or other stimuli that are predictive of an outcome (preferably a rewarding outcome, since 653 female odour is considered a positively arousing stimulus due to its effects on reproductive 654 behaviours).

655

In our study, mice clearly showed more interest in a filter paper infused with almond smell than in an odour-free filter paper. As expected, there was an observable trend of the overall exploration time to decrease over the course of the four trials, which is in line with the findings of Rattazzi, Cariboni, Poojara, Shoenfeld, & D'Acquisto (2015), who reported that mice habituated to the odours after the first exposure, since the odour exploration time was significantly reduced in the second and third exposures compared with the first exposure.

662

663 Rattazzi, Cariboni, Poojara, Shoenfeld, & D'Acquisto (2015) used a 100-fold dilution of almond 664 extract (10µl almond extract to 990µl distilled water) and the solution was prepared in the morning of 665 the same day of the test. In the present study, almond-odour infused filter papers and O^a objects were 666 also prepared on the day of each trial, but we did not use any dilutions (O^a objects had been labelled 667 with 100%-concentrated almond oil). Given that the potency of almond oil in our experiment was 668 likely higher than that used by above-mentioned authors, almond odour could have 'trumped' female 669 odour in animals' attention capture. This was, however, not the case - mice consistently showed 670 significantly higher exploration time and preference for O+ in Experiment 7. This observation was 671 very interesting, as it not only confirmed that mice are more attracted to female odour than to a non-672 social odour (which was already well-established in the literature), but it showed that this holds true, 673 even when almond odour is intense and found at a novel location. Experiment 7 represents a very 674 important addition to our study, because with it we were able to compare the degree of attention 675 allocation to female odour with that to another olfactory stimulus with lesser motivational 676 significance.

677

678 One explanation that could have accounted for the very high preference of mice for O+ compared 679 with O^a was that, in fact, the high concentration of almond oil resulted in the odour of O^a being 680 aversive to the mice. According to (Saraiva et al., 2016), some odours that are attractive at a low 681 concentration can become aversive in high concentrations. In our Experiment 7, however, we were 682 able to determine whether the mice were attracted or repulsed by the almond smell by looking at the 683 video recordings. Mice did not display avoidance behaviour toward O^a objects; they not only sniffed 684 the O^as multiple times upon their first discovery of such object in one of the Y-maze's arms, but they 685 also returned to this object several times during the 3-minute test. Considering that the Y-maze 686 contains three arms and that in Experiment 7, one arm was always object-free, and another arm 687 contained a highly attractive odour (O+), the animal could have easily avoided O^a by never returning 688 to the respective arm. Clearly, mice did not find O^as aversive, but they showed a distinct preference 689 for O+s.

690

691 In summary, the present study used object exploration measures to quantify the degree of attention 692 allocation by male mice to object novelty and/or female odour. Initial experiments demonstrated that 693 in the absence of other arousing features, objects with female odours capture more attention than an 694 odour-neutral object. These results agree with previous research showing that under laboratory 695 conditions, sexually isolated male mice are aroused by the odour of female conspecifics. The study 696 further demonstrated that odour 'wins' over location novelty in the degree of attention allocation and 697 that its motivational value is even greater in combination with location novelty. This supports the 698 conclusion that odour interacts with other arousing stimuli to form arousing contexts. Female odour 699 was able to capture more attention than a novel object, but only when combined with location novelty, 700 suggesting that the additive effects of visual and olfactory cues on attention exceed those of a single 701 strongly arousing stimulus. Finally, female odour attracted more attention than a mildly attractive 702 odour (low in motivational significance), suggesting that mice are able to form emotionally-charged 703 memories that are different from memories associated with other stimuli. These experiments 704 contribute to the understanding of the effects of female odour on value-modulated attention capture 705 and provide a reliable protocol to quantify attention allocation in mice. The findings obtained here 706 should encourage future research to use odour in investigating the influence of emotional arousal on 707 attention and memory.

- 708
- 709

- Acknowledgements We thank G. Winocur for an inspiring discussion, J. Neill for her support, and C. Charalambous for statistical advice.

713	References
714	
715	Aggleton, J.P., Albasser, M.M., Aggleton, D.J., Poirier, G.L., & Pearce, J.M. (2010). Lesions of the
716	rat perirhinal cortex spare the acquisition of a complex configural visual discrimination yet
717	impair object recognition. Behavioral Neuroscience, 124, 55-68. doi:10.1037/a0018320
718	Alkam, T., Hiramatsu, M., Mamiya, T., Aoyama, Y., Nitta, A., Yamada, K., Kim, H.C., &
719	Nabeshima, T. (2011). Evaluation of object-based attention in mice. Behavioural Brain
720	Research, 220, 185-193. doi:10.1016/j.bbr.2011.01.039
721	Anderson, B.A. (2016). Value-driven attentional capture in the auditory domain. Attention Perception
722	& Psychophysics, 78, 242-250. doi:10.3758/s13414-015-1001-7
723	Anderson, B.A., Laurent, P.A., & Yantis, S. (2011). Value-driven attentional capture. Proceedings of
724	the National Academy of Sciences of the United States of America, 108, 10367-10371.
725	doi:10.1073/pnas.1104047108
726	Anderson, B.A., Laurent, P.A., & Yantis, S. (2014). Value-driven attentional priority signals in
727	human basal ganglia and visual cortex. Brain Research, 1587, 88-96.
728	doi:10.1016/j.brainres.2014.08.062
729	Antunes, M., & Biala, G. (2012) The novel object recognition memory: Neurobiology, test procedure,
730	and its modifications. Cognitive Processing, 13, 93-110. doi:10.1007/s10339-011-0430-z
731	Arbuckle, E. P., Smith, G. D., Gomez, M. C., & Lugo, J. N. (2015). Testing for Odor Discrimination
732	and Habituation in Mice. Jove-Journal of Visualized Experiments(99), 7. doi:10.3791/52615
733	Aston-Jones, G., Rajkowski, J., Kubiak, P., Valentino, R.J., & Shipley, M.T. (1996). Role of the locus
734	coeruleus in emotional activation. Emotional Motor System, 107, 379-402. Retrieved from
735	https://doi.org/10.1016/S0079-6123(08)61877-4
736	Barker, G.R.I., & Warburton, E.C. (2011). When is the hippocampus involved in recognition
737	memory?. Journal of Neuroscience, 31, 10721-10731. doi:10.1523/JNEUROSCI.6413-10.2011
738	Beny, Y., & Kimchi, T. (2014). Innate and learned aspects of pheromone-mediated social behaviours.
739	Animal Behaviour, 97, 301-311. doi:10.1016/j.anbehav.2014.09.014
740	Berlyne, D.E. (1950). Novelty and curiosity as determinants of exploratory behaviour. British
741	Journal of Psychology, 41, 68-80. doi:10.1111/j.2044-8295.1950.tb00262.x
742	Brennan, P.A., & Kendrick, K.M. (2006). Mammalian social odours: Attraction and individual
743	recognition. Philosophical Transactions of the Royal Society B-Biological Sciences, 361, 2061-
744	2078. doi:10.1098/rstb.2006.1931
745	Burke, S.N., Wallace, J.L., Nematollahi, S., Uprety, A.R., & Barnes, C.A. (2010). Pattern Separation
746	Deficits May Contribute to Age-Associated Recognition Impairments. Behavioral
747	Neuroscience, 124, 559-573. doi:10.1037/a0020893

- Chelazzi, L., Estocinova, J., Calletti, R., Lo Gerfo, E., Sani, I., Della Libera, C., & Santandrea, E.
 (2014). Altering spatial priority maps via reward-based learning. *Journal of Neuroscience*, *34*, 8594-8604. doi:10.1523/JNEUROSCI.0277-14.2014
- 751 Chun, M.M., Golomb, J.D., & Turk-Browne, N.B. (2011). A taxonomy of external and internal
 752 attention. *Annual Review of Psychology*, 62, 73-101. doi:
 753 10.1146/annurev.psych.093008.100427
- Compton, R.J., 2003. The interface between emotion and attention: A review of evidence from
 psychology and neuroscience. *Behavioral and Cognitive Neuroscience Reviews*, 2, 115–129.
 doi:10.1177/1534582303255278
- 757 Connor, J. (1972). Olfactory control of aggressive and sexual behavior in mouse (Mus-musculus-L.).
 758 *Psychonomic Science*, 27, 1-3. doi:10.3758/BF03328867
- Corbetta, M., Patel, G., & Shulman, G.L. (2008). The reorienting system of the human brain: From
 environment to theory of mind. *Neuron*, *58*, 306-324. doi:10.1016/j.neuron.2008.04.017
- 761 Corbetta, M., & Shulman, G.L. (2002). Control of goal-directed and stimulus-driven attention in the
 762 brain. *Nature Reviews Neuroscience*, *3*, 201-215. doi:10.1038/nrn755
- Davis, K.E., Eacott, M.J., Easton, A., & Gigg, J. (2013). Episodic-like memory is sensitive to both
 Alzheimer's-like pathological accumulation and normal ageing processes in mice. *Behavioural Brain Research*, 254, 73-82. doi:10.1016/j.bbr.2013.03.009
- Desimone, R., & Duncan, J. (1995). Neural mechanisms of selective visual-attention. *Annual Review of Neuroscience*, 18, 193-222. doi:10.1146/annurev.ne.18.030195.001205
- Devore, S., Lee, J., & Linster, C. (2013). Odor Preferences Shape Discrimination Learning in Rats. *Behavioral Neuroscience*, *127*, 498-504. doi:10.1037/a0033329
- Duffaud, A.M., Killcross, S., & George, D.N. (2007). Optional-shift behaviour in rats: A novel
 procedure for assessing attentional processes in discrimination learning. *Quarterly Journal of Experimental Psychology*, 60, 534-542. doi:10.1080/17470210601154487
- Eacott, M.J., & Norman, G. (2004). Integrated memory for object, place, and context in rats: A
 possible model of episodic-like memory?. *Journal of Neuroscience*, 24, 1948-1953.
 doi:10.1523/jneurosci.2975-03.2004
- 776 Ennaceur, A. (2010). One-trial object recognition in rats and mice: Methodological and theoretical
 777 issues. *Behavioural Brain Research*, *215*, 244-254. doi:10.1016/j.bbr.2009.12.036
- Ennaceur, A., Neave, N., & Aggleton, J.P. (1997). Spontaneous object recognition and object location
 memory in rats: The effects of lesions in the cingulate cortices, the medial prefrontal cortex, the
 cingulum bundle and the fornix. *Experimental Brain Research*, 113, 509-519.
 doi:10.1007/pl00005603
- Esber, G.R., & Haselgrove, M. (2011). Reconciling the influence of predictiveness and uncertainty on
 stimulus salience: A model of attention in associative learning. *Proceedings of the Royal Society B-Biological Sciences*, 278, 2553-2561. doi:10.1098/rspb.2011.0836

- Failing, M.F., & Theeuwes, J. (2014). Exogenous visual orienting by reward. *Journal of Vision, 14*, 19. doi:10.1167/14.5.6
- Foley, N.C., Jangraw, D.C., Peck, C., & Gottlieb, J. (2014). Novelty enhances visual salience
 independently of reward in the parietal lobe. *Journal of Neuroscience*, 34, 7947-7957.
 doi:10.1523/jneurosci.4171-13.2014
- Gaskin, S., Tardif, M., Cole, E., Piterkin, P., Kayello, L., & Mumby, D.G. (2010). Object
 familiarization and novel-object preference in rats. *Behavioural Processes*, *83*, 61-71.
 doi:10.1016/j.beproc.2009.10.003
- George, D.N., & Pearce, J.M. (1999). Acquired distinctiveness is controlled by stimulus relevance not
 correlation with reward. *Journal of Experimental Psychology-Animal Behavior Processes*, 25,
 363-373. doi:10.1037/0097-7403.25.3.363
- 796 Gitelman, D.R. (2003). Attention and its disorders. *British Medical Bulletin*, 65, 21-34.
 797 doi:10.1093/bmb/65.1.21
- Golomb, J.D., Nguyen-Phuc, A.Y., Mazer, J.A., McCarthy, G., & Chun, M.M. (2010). Attentional
 facilitation throughout human visual cortex lingers in retinotopic coordinates after eye
 movements. *Journal of Neuroscience*, *30*, 10493-10506. doi:10.1523/jneurosci.1546-10.2010
- Hartikainen, M.K., Ogawa, H.K., Soltani, T.M., & Knight, T.R. (2007). Emotionally arousing stimuli
 compete for attention with left hemispace. *NeuroReport*, 18, 1929-1933.
 doi:10.1097/WNR.0b013e3282f1ca18
- Horvitz, J.C. (2000). Mesolimbocortical and nigrostriatal dopamine responses to salient non-reward
 events. *Neuroscience*, *96*, 651–656. doi:10.1016/S0306-4522(00)00019-1
- Huckins, L. M., Logan, D. W., & Sanchez-Andrade, G. (2013). Olfaction and olfactory-mediated
 behaviour in psychiatric disease models. *Cell and Tissue Research*, 354, 69-80.
 doi:10.1007/s00441-013-1617-7
- Johnston, R.E. (2003). Chemical communication in rodents: From pheromones to individual
 recognition. *Journal of Mammalogy*, *84*, 1141-1162. doi:10.1644/BLe-010
- Kadohisa, M. (2013). Effects of odor on emotion, with implications. *Frontiers in Systems Neuroscience*, 7, 66. doi:10.3389/fnsys.2013.00066
- Kakade, S., & Dayan, P. (2002). Dopamine: generalization and bonuses. *Neural Networks*, 15, 549 –
 559. doi:10.1016/S0893-6080(02)00048-5
- Kalish, M. L., & Kruschke, J. K. (2000). The role of attention shifts in the categorization of
 continuous dimensioned stimuli. *Psychological Research-Psychologische Forschung*, 64, 105–
 116. doi:10.1007/s004260000028
- Katsuki, F., & Constantinidis, C. (2014). Bottom-up and top-down attention: Different processes and
 overlapping neural systems. *Neuroscientist*, *20*, 509-521. doi:10.1177/1073858413514136

- Kaye, H., & Pearce, J.M. (1984). The strength of the orienting response during Pavlovian
 conditioning. *Journal of Experimental Psychology-Animal Behavior Processes*, 10, 90-109.
 doi:10.1037/0097-7403.10.1.90
- Laurent, P.A. (2008). The emergence of saliency and novelty responses from reinforcement learning
 principles. *Neural Networks*, 21, 1493–1499. doi:10.1016/j.neunet.2008.09.004
- Lawrence, A. D., Sahakian, B. J., Rogers, R. D., Hodges, J. R., & Robbins, T. W. (1999).
 Discrimination, reversal, and shift learning in Huntington's disease: Mechanisms of impaired
 response selection. *Neuropsychologia*, *37*, 1359–1374. doi:10.1016/S0028-3932(99) 00035-4
- Le Pelley, M.E., Mitchell, C.J., Beesley, T., George, D.N., & Wills, A.J. (2016). Attention and
 associative learning in humans: An integrative review. *Psychological Bulletin, 142*, 1111-1140.
 doi:10.1037/bul0000064
- Lee, T.-H., Sakaki, M., Cheng, R., Velasco, R., & Mather, M. (2014). Emotional arousal amplifies the
 effects of biased competition in the brain. *Social Cognitive and Affective Neuroscience*, 9,
 2067-2077. doi:10.1093/scan/nsu015
- Levine, L.J., & Edelstein, R.S. (2009). Emotion and memory narrowing: A review and goal-relevance
 approach. *Cognition & Emotion*, 23, 833-875. doi:10.1080/02699930902738863
- 836 Mackintosh, J.H. (1970). Territory formation by laboratory mice. *Animal Behaviour, 18*, 177-183.
 837 doi:10.1016/0003-3472(70)90088-6
- Mackintosh, N. J. (1975). A theory of attention: Variations in the associability of stimuli with
 reinforcement. *Psychological Review*, *82*, 276–298. doi:10.1037/h0076778
- McGaughy, J., & Sarter, M. (1998). Sustained attention performance in rats with intracortical infusions of 192 IgG-saporin-induced cortical cholinergic deafferentation: Effects of physostigmine and FG 7142. *Behavioral Neuroscience*, *112*, 1519-1525. doi:10.1037/07357044.112.6.1519
- Michael, G.A., Jacquot, L., Millot, J.L., & Brand, G. (2003). Ambient odors modulate visual
 attentional capture. *Neuroscience Letters*, *352*, 221-225. doi:10.1016/s0304-3940(03)01078-4
- 846 Miller, E.K., & Cohen, J.D. (2001). An integrative theory of prefrontal cortex function. *Annual*847 *Review of Neuroscience, 24*, 167-202. doi:10.1146/annurev.neuro.24.1.167
- Mumby, D.G., Glenn, M.J., Nesbitt, C., & Kyriazis, D.A. (2002). Dissociation in retrograde memory
 for object discriminations and object recognition in rats with perirhinal cortex damage. *Behavioural Brain Research*, 132, 215-226. doi:10.1016/s0166-4328(01)00444-2
- Murai, T., Okuda, S., Tanaka, T., & Ohta, H. (2007). Characteristics of object location memory in
 mice: Behavioral and pharmacological studies. *Physiology & Behavior*, 90, 116-124.
 doi:10.1016/j.physbeh.2006.09.013
- Ohman, A., & Mineka, S. (2001). Fears, phobias, and preparedness: Toward an evolved module of
 fear and fear learning. *Psychological Review*, *108*, 483-522. doi:10.1037//0033-295x.108.3.483

- Oliveira, A.M.M., Hawk, J.D., Abel, T., & Havekes, R. (2010). Post-training reversible inactivation
 of the hippocampus enhances novel object recognition memory. *Learning & Memory*, 17, 155160. doi:10.1101/lm.1625310
- Pantoja, J., Ribeiro, S., Wiest, M., Soares, E., Gervasoni, D., Lemos, N.A.M., & Nicolelis, M.A.L.
 (2007). Neuronal activity in the primary somatosensory thalamocortical loop is modulated by
 reward contingency during tactile discrimination. *Journal of Neuroscience*, *27*, 10608-10620.
 doi:10.1523/JNEUROSCI.5279-06.2007
- Pearce, J.M., & Hall, G. (1980). A model for Pavlovian learning: Variations in the effectiveness of
 conditioned but not of unconditioned stimuli. *Psychological Review*, 87, 532-552.
 doi:10.1037/0033-295x.87.6.532
- Pessoa, L. (2009). How do emotion and motivation direct executive control?. *Trends in Cognitive Sciences*, 13, 160-166. doi:10.1016/j.tics.2009.01.006
- Piper, B.J., Fraiman, J.B., & Meyer, J.S. (2005). Repeated MDMA ("Ecstasy") exposure in adolescent
 male rats alters temperature regulation, spontaneous motor activity, attention, and serotonin
 transporter binding. *Developmental Psychobiology*, 47, 145-157. doi:10.1002/dev.20085
- Pourtois, G., Schettino, A., & Vuilleumier, P. (2013). Brain mechanisms for emotional influences on
 perception and attention: What is magic and what is not. *Biological Psychology*, *92*, 492-512.
 doi: 10.1016/j.biopsycho.2012.02.007
- 874 Pratt, J., & Hommel, B. (2003). Symbolic control of visual attention: The role of working memory
 875 and attentional control settings. *Journal of Experimental Psychology-Human Perception and*876 *Performance, 29*, 835-845. doi:10.1037/0096-1523.29.5.835
- Qi, S.Q., Zeng, Q.H., Ding, C., & Li, H. (2013). Neural correlates of reward-driven attentional
 capture in visual search. *Brain Research*, *1532*, 32-43. doi:10.1016/j.brainres.2013.07.044
- Rattazzi, L., Cariboni, A., Poojara, R., Shoenfeld, Y., & D'Acquisto, F. (2015). Impaired sense of
 smell and altered olfactory system in RAG-1(-/-) immunodeficient mice. *Frontiers in Neuroscience*, 9, 9. doi:10.3389/fnins.2015.00318
- Saraiva, L. R., Kondoh, K., Ye, X. L., Yoon, K. H., Hernandez, M., & Buck, L. B. (2016).
 Combinatorial effects of odorants on mouse behavior. *Proceedings of the National Academy of Sciences of the United States of America*, 113, E3300-E3306. doi:10.1073/pnas.1605973113
- Sarter, M., Givens, B., & Bruno, J.P. (2001). The cognitive neuroscience of sustained attention:
 Where top-down meets bottom-up. *Brain Research Reviews*, *35*, 146-160. doi:10.1016/s01650173(01)00044-3
- Shapiro, K.L., Egerman, B., & Klein, R.M. (1984). Effects of arousal on human visual dominance. *Perception & Psychophysics*, 35, 547-552. doi:10.3758/bf03205951
- Silvers, J.M., Harrod, S.B., Mactutus, C.F., & Booze, R.M. (2007). Automation of the novel object
 recognition task for use in adolescent rats. *Journal of Neuroscience Methods*, *166*, 99-103. doi:
 10.1016/j.jneumeth.2007.06.032

- Spence, C. (2010). Crossmodal spatial attention. *Annals of The New York Academy of Sciences*, 1191,
 182-200. doi:10.1111/j.1749-6632.2010.05440.x
- 895 Sutherland, N. S., & Mackintosh, N. J. (1971). *Mechanisms of animal discrimination learning*. New
 896 York, NY: Academic Press
- Talmi, D. (2013). Enhanced emotional memory: cognitive and neural mechanisms. *Current Directions in Psychological Science*, 22, 430-436. doi:10.1177/0963721413498893
- Talmi, D., & McGarry, L.M. (2012). Accounting for immediate emotional memory enhancement.
 Journal of Memory and Language, 66, 93-108. doi:10.1016/j.jml.2011.07.009
- 901 Talmi, D., Slapkova, M., & Wieser, M. (2018). Testing the possibility of model-based Pavlovian
 902 control on attention to threat. *Journal of Cognitive Neuroscience*, 31, 36-48. doi:
 903 10.1162/jocn a 01329
- 904 Vermeulen, N., Godefroid, J., & Mermillod, M. (2009). Emotional modulation of attention: Fear 905 increases but disgust reduces the attentional blink. Plos One. 4. 5. 906 doi:10.1371/journal.pone.0007924
- 907 Vuilleumier, P. (2005). How brains beware: Neural mechanisms of emotional attention. *Trends in* 908 *Cognitive Sciences*, 9, 585-594. doi:10.1016/j.tics.2005.10.011
- Wang, L.H., Duan, Y.Y., Theeuwes, J., & Zhou, X.L. (2014). Reward breaks through the inhibitory
 region around attentional focus. *Journal of Vision*, 14, 7. doi:10.1167/14.12.2
- Wilson, P.N., Boumphrey, P., & Pearce, J.M. (1992). Restoration of the orienting response to a light
 by a change in its predictive accuracy. *Quarterly Journal of Experimental Psychology Section B-Comparative and Physiological Psychology*, 44, 17-36. doi:10.1080/02724999208250600
- 914 Yang, M., & Crawley, J. N. (2009). Simple behavioral assessment of mouse olfaction. *Current* 915 protocols in neuroscience, Chapter 8, Unit 8.24, doi: 10.1002/0471142301.ns0824s48
- 916 Yantis, S., Schwarzbach, J., Serences, J.T., Carlson, R.L., Steinmetz, M.A., Pekar, J.J., & Courtney,
 917 S.M. (2002). Transient neural activity in human parietal cortex during spatial attention shifts.
 918 Nature Neuroscience, 5, 995-1002. doi:10.1038/nn921
- Zaborszky, L., Gaykema, R.P., Swanson, D.J., & Cullinan, W.E. (1997). Cortical input to the basal
 forebrain. *Neuroscience*, *79*, 1051-1078. doi:10.1016/s0306-4522(97)00049-3
- 921 Wentura, D., Muller, P., & Rothermund, K. (2014). Attentional capture by evaluative stimuli: Gain922 and loss-connoting colors boost the additional- singleton effect. *Psychonomic Bulletin &*923 *Review*, 21, 701–707. doi:10.3758/s13423-013-0531-z
- 924
- 925