1	Sex differences in fear regulation and reward seeking behaviors in a fear-safety-reward		
2	discrimination task		
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5	Abbreviated title: Sex differences in conditioned safety		
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28 Abstract

- 29 Reward availability and the potential for danger or safety potently regulates emotion. Despite
- 30 women being more likely than men to develop emotion dysregulation disorders, there are
- 31 comparatively few studies investigating fear, safety and reward regulation in females. Here, we
- 32 show that female Long Evans rats do not suppress conditioned freezing in the presence of a
- 33 safety cue, nor do they extinguish their freezing response, whereas males do both. Females
- 34 were also more reward responsive during the reward cue until the first footshock exposure, at
- 35 which point there were no sex differences in reward seeking to the reward cue. Darting analyses
- 36 indicate females might be able to regulate this behavior in response to the safety cue,
- 37 suggesting they might be able to discriminate between fear and safety cues but do not
- 38 demonstrate this with conditioned suppression of the freezing behavior. However, levels of
- 39 darting in this study were too low to make any clear conclusions. In summary, females showed
- 40 a significantly different behavioral profile than males in a task that tests the ability to discriminate
- 41 among fear, safety and reward cues. This paradigm offers a great opportunity to test for
- 42 mechanisms that are generating these behavioral sex differences in learned safety and reward
- 43 seeking.
- 44
- 45 Keywords: sex differences; fear; safety; reward

46 **1. Introduction**

47 Clinical disorders arising from maladaptive emotion regulation present a large burden on society 48 worldwide. Many of these disorders show comorbidity, for example, addiction with anxiety 49 disorders (Grant et al., 2016). Cues predicting something aversive elicit avoidance and fear 50 behaviors whereas cues predicting reward elicit approach and reward-seeking behaviors. Cues 51 signifying safety have the power to modulate fear and reward-seeking behaviors by informing 52 the organism whether or not the environment is safe (Walasek, Wesierska, & Zieliński, 1995). 53 Thus, safety, fear and reward behaviors, and the circuitries governing these behaviors, are 54 intertwined. The majority of studies on reward and fear processing have been conducted in 55 parallel, investigating the circuitries separately in primarily male subjects. If we hope to 56 understand and treat comorbid disorders resulting from maladaptive emotion regulation, 57 increased efforts in investigating how these circuitries integrate their functions to influence 58 behavior is needed in both male and female subjects. 59 60 Our laboratory has designed and validated a behavioral task in which fear, safety and reward 61 cues are learned within the same session allowing us to assess the animal's ability to 62 discriminate among these cues (Ng et al., 2018; Sangha et al., 2013; Sangha, Greba et al., 63 2014; Sangha, Robinson et al., 2014). Rats are exposed to cues associated with safety, fear 64 (fear cue paired with footshock), and reward (reward cue paired with sucrose). Male rats 65 consistently learn to discriminate among safety, fear and reward cues to 1) suppress 66 conditioned freezing in the presence of a safety cue (fear+safety cue), and 2) increase reward 67 seeking when reward is available (reward cue) (Ng et al., 2018; Sangha et al., 2013; Sangha, 68 Greba, et al., 2014; Sangha, Robinson, et al., 2014). This paradigm also allows us to investigate

69 how safety cues can regulate both fear and reward behaviors. Evidence suggests that reward

70 learning mechanisms overlap with safety learning (Leknes et al., 2011; Pollak et al., 2008;

Rescorla, 1969; Rogan et al., 2005; Sangha et al., 2013; Tanimoto et al., 2004; Walasek et al.,

1995). For example, learned safety can act as a behavioral antidepressant in mice (Pollak et al.,

73 2008), and animals will perform certain behaviors in order to turn on a safety signal (Rescorla,

1969; Rogan et al., 2005). Within the amygdala we have shown a subpopulation of neurons

responding with the same level of excitation or inhibition during both the reward and safety cues

76 (Sangha et al., 2013). Additionally, we have shown inactivation of the prelimbic or infralimbic

77 cortices of the ventromedial prefrontal cortex have differential effects on reward and safety

discrimination, respectively (Sangha, Robinson, et al., 2014). Thus, in male rats, our prior work

has already shown a critical involvement of the corticoamygdalar circuit in learning this fearsafety-reward cue discrimination.

81

82 Much of the research investigating emotion regulation mechanisms have exclusively used male 83 subjects. In a study using male Vietnam veterans, PTSD patients show impairments in 84 suppressing their fear response in the presence of a safety cue (Jovanovic et al., 2009). But, 85 women are more than twice as likely to develop Post-Traumatic Stress Disorder (PTSD) than 86 men, with females having a lifetime prevalence of 8.5% in contrast to 3.4% in males (Mclean et 87 al., 2011). In fear studies that have included female rats, it has been shown that females exhibit 88 lower levels of freezing behavior than male rats after repeated fear cue presentations (Daviu et 89 al., 2014). These findings have been thought to indicate a difficulty in fear conditioning learning 90 in female rats. A more recent experiment has identified that approximately 40% of female rats 91 tested exhibit an alternate fear behavior in the form of fast paced movements called 'darting': 92 this was only seen in approximately 10% of male rats tested (Gruene et al., 2015). There is also 93 evidence of sex differences in the seeking of natural rewards, where it has been reported that 94 female rats consume more sucrose pellets than males and are willing to work harder for them 95 (Tapia, Lee, Weise, Tamasi, & Will, 2019). Dopamine signaling during reward tasks has also 96 been demonstrated to be different between sexes. For example, Conway et al (2019) showed 97 females continue to perform intracranial self-stimulation for brain stimulation reward while under 98 the influence of a kappa-opioid receptor agonist, which suppresses dopamine release, whereas 99 males decrease this behavior. Their data suggest that female rats may have an increased 100 capacity to produce and release dopamine compared to males, under these conditions. Our 101 prior work has shown, in males, that dopamine signaling in the basolateral amygdala contributes 102 to effective discrimination among fear, safety and reward cues (Ng et al., 2018). 103

Taken together, we hypothesized there would be sex differences in the ability to express clear discrimination among fear, safety and reward cues. The inability of male PTSD patients to learn safety signaling has been labeled a biomarker of the disorder (Jovanovic et al., 2012). Due to sex-related differences in human diagnosis of PTSD, with women diagnosed at rates twice that of men (Glover et al., 2015), any differences female rats have in the learning or retention of safety signals could steer towards further research on the neurological processes underlying these variations.

111 **2. Materials and Methods**

112 2.1 Subjects

113 A total of 24 adult male (215-375g) and 28 adult age-matched female (198-230g) Long Evans

114 rats (Blue Spruce; Envigo, Indianapolis), were single-housed and handled for 1 week prior to

115 testing. All procedures were performed during the light cycle and approved by the Purdue

- 116 Animal Care and Use Committee. Rats had *ad libitum* access to food and water prior to the
- 117 start of the experiment. After experiment onset, they were maintained on a food restricted diet
- 118 (20g per day for males; 16g per day for females) until the last day of the experiment.
- 119

120 2.2 Apparatus

121 The rats were trained in operant conditioning chambers consisting of Plexiglas boxes (32cm

122 length x 25cm width x 30cm height) encased in sound-attenuating chambers (Med Associates,

123 ST Albans, VT). 10% liquid sucrose was delivered through a recessed port 2cm above the floor

124 in the center of one wall. Two lights (28V, 100mA) were located 10.5cm from floor on either side

125 of the port. A light (28V, 100mA) 27cm above the floor on the wall was on throughout the entire

126 session. Auditory cues were delivered via a speaker (ENV-224BM) located 24cm from the floor

127 on the same wall as the port. Footshocks were delivered through a grid floor via a constant

128 current aversive stimulator (ENV-414S). An overhead video camera and side video camera

129 recorded the sessions for subsequent offline video scoring.

130

131 2.3 Behavioral Procedures

132 <u>Reward pre-training</u> (5 sessions): An auditory cue is paired with 10% sucrose solution delivery
 133 (100µl) and serves as the reward cue (25 trials; ITI, 90-130s).

134 Habituation (1 session): Rats continue to receive 25 reward cue-sucrose pairings (ITI, 90-130s)

135 in addition to 5 unreinforced presentations each of the future fear and safety cues in order to

136 habituate the rats to their presentation, thereby reducing any baseline freezing to these novel

137 cues.

138 Discriminative conditioning (DC) (4 sessions): Reward cue-sucrose pairings continue (15 trials).

139 Another auditory cue is paired with a mild 0.5mA, 0.5s footshock and serves as the fear cue (4

trials). In separate trials the 20s fear cue is presented at the same time as a 20s safety light cue

141 resulting in no footshock ('fear+safety', 15 trials). Trials in which the safety cue is presented

142 alone without any footshock are also included to assess whether freezing develops to the safety

143 cue as well as providing the animal with additional trials that contains a safety cue-no shock

144 contingency (10 trials). Trials are presented pseudorandomly (ITI, 100-140 s). Cues were

145 counterbalanced across subjects with the caveat that the fear+safety compound cue was 146 composed of one auditory cue and one light cue. Eight of the male rats and 12 of the female 147 rats underwent DC training in which the reward cue was a continuous auditory cue (3 kHz, 20s 148 cue; 70dB), the fear cue a pulsing auditory cue (11 kHz, 20s; 70dB), and the safety cue was the 149 presentation of two lights (28V, 100mA located on both sides of the port). The remaining eight 150 male rats and eight female rats underwent training in which the fear and safety cue stimuli were 151 counterbalanced: the light served as the fear cue and the pulsing auditory cue served as the 152 safety cue. 153 Extinction Training (1 session): One day after the last DC session, both the reward cue and fear

154 cue were presented 20 times each in a pseudorandomized order without sucrose or footshock.

155 <u>Extinction Test (1 session)</u>: One day after extinction training, rats were presented with the

reward (10 trials), fear (10 trials), fear+safety (5 trials) and safety (5 trials) cues in a

157 pseudorandomized order (ITI, 60-120s). None of the cues were presented with sucrose or

- 158 footshock.
- 159

160 To exclude possible sex differences in pain sensitivity and foot shock perception a separate

161 group of male (n=8) and age-matched female (n=8) rats was presented with a series of

unsignalled foot shocks of increasing intensities (0.3 mA, 0.35 mA, 0.4 mA, 0.45 mA, 0.5 mA,

163 0.55 mA, 0.6 mA, 0.7 mA, 0.8 mA, 0.9, 1.0 mA) with an inter-stimulus interval of 2 min. The

164 session was flanked with 5 min intervals in which no stimulus occurred.

165

166 2.4 Data analyses

167 Our experimental groups to directly compare males and females on discrimination behavior 168 consisted of 16-20 rats. Cohorts of 4 or 8 female rats were trained alongside cohorts of 4 male 169 rats for a total of 4 replications. Fear behavior was assessed manually offline from videos by 170 measuring freezing, defined as complete immobility with the exception of respiratory 171 movements, which is an innate defensive behavior (Blanchard & Blanchard, 1969; Fendt & 172 Fanselow, 1999). The total time spent freezing during each 20s cue was quantified and 173 expressed as a percentage. Measuring the total time the animal spent inside the reward port 174 and at the entrance of the port with nose positioned at port entrance during each cue assessed 175 reward seeking behavior and was expressed as a percentage. Darting behavior was detected 176 and quantified offline from videos recorded from overhead cameras via a custom MatLab 177 program, with movements of a velocity of 23.5cm/s or faster gualifying as a single dart (Gruene 178 et al., 2015); these were also confirmed manually. Darting was expressed as the averaged # of 179 darts per cue (sum of darts/ # trials) or trial (sum of darts). Since there were different number of 180 trials per reward, fear, fear+safety and safety cue in each DC session and test for extinction, 181 this was expressed as the sum of darts across trials divided by the number of trials for each cue 182 (sum of darts/ # trials). During extinction training, data for each individual trial is shown and the 183 dart rate is expressed the averaged sum of darts for each individual trial (sum of darts). Three 184 individuals performed manual offline behavioral scoring. Pearson's correlations of behavioral 185 values between scorers were greater than r = 0.80. Behavioral data were analyzed with one-186 way or two-way repeated measures ANOVAs, with sex as the independent factor and condition 187 as the repeated factor, followed by *post hoc* Sidak's, Tukey's or Dunnett's multiple comparisons 188 tests with GraphPad Prism 8. P values were adjusted for multiple comparisons. 189

190 For shock sensitivity testing, freezing duration in the 2-min intervals between shock

191 presentations was scored manually as an indicator of fear, as well as darting and jumping as an

192 immediate shock response. For the freezing durations, a two-way repeated measures ANOVA

193 was carried out via GraphPad Prism 7, with sex as the independent factor and shock intensity

194 as the repeated factor. Darting and jumping were assessed as dichotomous variables with

195 darting/no darting and jumping/no jumping, respectively. For both, a Cochran test was

196 performed.

197

198 **3. Results**

199 3.1 Female rats spent more time reward seeking during reward pre-training

200 All rats first underwent 5 reward pre-training sessions in which the reward cue was paired with

201 sucrose delivery. The percent time spent at or in the reward port during each reward cue across

202 each reward session was quantified (Figure 1b). Two-way repeated-measures ANOVAs showed

203 main effects of session (F(4,136)=5.395, p=0.0005) and sex (F(1,34)=10.83, p=.0023), but no

significant interaction (F(4,136)=0.9031, p=0.4641). *Post hoc* Sidak's multiple comparisons test

205 showed females spent significantly more time reward seeking during the reward cue than males

- for sessions R2 (p=0.0274), R3 (p=0.0151) and R5 (p=0.0041). The latency, in seconds, to
- 207 enter the port post-cue onset was also calculated for each reward cue presentation across all
- 208 sessions (Figure 1c). Two-way repeated-measures ANOVAs showed a main effect of sex
- 209 (F(1,34)=20.37, p<.0001), but no significant interaction (F(4,136)=1.684, p=0.1571) or main
- effect of session (F(4,136)=0.7755, p=0.5429). *Post hoc* Sidak's multiple comparisons test
- showed females were significantly faster to enter the port than males during the last 3 reward
- sessions (R3, p=0.001; R4, p=0.0391; R5, p=0.0014). Taken together, female rats consistently

spent more time than males in the reward port during the reward cue in reward pre-training

- 214 sessions.
- 215

216 3.2 Female rats did not show conditioned inhibition of freezing

217 After reward pre-training, rats were then exposed to sessions also consisting of reward, fear and

218 safety cues. The reward cue and sucrose reward were the same as the reward pre-training

sessions. The fear cue was paired with a 0.5mA footshock, and both the safety cue and

220 fear+safety cue did not result in footshock or sucrose.

221

222 The percent time spent at or in the reward port during each cue across session was quantified 223 for each DC session (Figure 2b). Two-way repeated-measures ANOVAs showed a significant 224 cue by sex effect, as well as main effects of cue and sex for DC1 (Table 1). Post hoc Sidak's 225 multiple comparisons test showed that, during DC1, females spent significantly more time 226 reward seeking during the reward cue compared to males (p<0.001), consistent to what was 227 seen in reward pre-training. For the remaining DC2-4 sessions, a main effect of cue was 228 observed (Table 1) and post hoc Sidak's multiple comparisons test showed that both male and 229 female rats spent significantly more time reward seeking during the reward cue compared to all 230 other cues (p<0.0001), with no significant differences between the males and females. Thus, 231 the noticeable increase in reward seeking in the females, that was seen during reward pretraining, dissipated by the 2nd DC session. 232

233

234 The percent time freezing during each cue across session was quantified for each DC session

235 (Figure 2c). Two-way repeated-measures ANOVAs showed a significant cue by sex effect for

236 sessions DC2-4, as well as main effects of cue and sex for every session (Table 1). *Post hoc*

- 237 Sidak's multiple comparisons tests showed that, for every session, females displayed
- significantly more freezing to the fear+safety cue compared to males (DC1, p=0.0313; DC2,

p=0.007; DC3, p=0.0007; DC4, p<0.0001). Females also showed significantly higher freezing

240 levels to the fear cue compared to males during DC2 (p=0.0111). Males showed a significant

reduction in freezing levels to the fear+safety cue compared to the fear cue during sessions

DC3 (p=0.0156) and DC4 (p<0.0001), thus showing significant conditioned inhibition of freezing.

243 Females did not show a significant inhibition of freezing during any session.

244

The number of darts during each cue was also quantified for each DC session and expressed
as a dart rate (Figure 2d; sum of darts/ # trials). Darting behavior during cue presentation was

- largely absent until DC3 and DC4. Two-way repeated-measures ANOVAs showed a significant
 cue by sex effect for DC4, as well as main effects of cue, for DC2-4, and sex, for DC1 and DC4
 (Table 1). *Post hoc* Sidak's multiple comparisons test showed that, during DC4, females
- 250 expressed more darting behavior compared to males during both the fear cue (p=0.0017) and
- the fear+safety cue (p=0.0186). The females reduced their darting rate from 0.51 during the fear
- cue to 0.32 during the fear+safety cue, but this was not statistically significant.
- 253
- 254 3.3 Female rats did not show significant extinction of freezing
- 255 The day after the last DC session all rats underwent fear and reward extinction within the same
- session. That is, both the fear and reward cues were presented within the same training
- 257 session, without footshocks or sucrose presentations.
- 258
- During extinction of reward, there was no main effect of reward trial (F(19,646)=1.526,
- 260 p=0.0704) or sex (F(1,34)=1.31, p=0.2603) and no interaction (F(19,646)=0.8927, p=0.5924);
- there was also no significant difference between male and female groups for any trial (Figure
- 262 3Bi). One day later when rats were re-tested for extinction memory (Figure 3Bii), there was a
- 263 main effect of cue (2-way RM ANOVA; F(3,102)=134.7, p<0.0001) and sex (F(1,34)=6.217,
- 264 p=0.0177). *Post hoc* Sidak's multiple comparisons test showed females had significantly more
- 265 port activity than males just during the safety cue (p=0.0452), although this difference did not
- reflect a large increase in port activity as females spent 6.38% +/- 0.86 of the safety cue in the
- 267 port compared to 2.66% +/- 0.86 in males. Overall, there appeared to be no differences in the
- ability of males and females to extinguish their reward seeking responses.
- 269

270 To assess fear extinction the averaged percent time freezing during each trial of fear extinction

- training was calculated (Figure 3Ci). There was a main effect of fear trial (2-way RM ANOVA;
- 272 F(19, 646)=7.69, p<0.0001) and sex (2-way RM ANOVA; F(1, 34)=4.607, p=0.0391), but no
- significant interaction (F(19, 646)=1.566, p=0.059). Compared to trial 1, males showed
- significantly reduced freezing in extinction trials 8, 9 and 11-20 (post hoc Tukey's multiple
- 275 comparisons test, p<0.05), demonstrating good fear extinction beginning around the 8th trial. In
- 276 contrast, females only showed a significant reduction in freezing during trial 19 compared to the
- first trial (*post hoc* Tukey's multiple comparisons test, p=0.0394), demonstrating relatively
- absent fear extinction. One day later when rats were retested for extinction memory (Figure
- 279 3Cii), there was a main effect of cue (F(3,102)=134.7, p<0.0001) and sex (F(1, 34)=6.217,
- 280 p=0.0177), as well as a significant interaction of cue X sex (F(3, 102)=3.481, p=0.0187). Post

hoc Sidak's multiple comparisons test showed that females froze significantly more than males
 to the fear (p=0.0146) and fear+safety (p=0.0091) cues. This indicates the continued absence of
 any extinction of freezing in females.

284

285 In response to each fear cue presentation across extinction, we also assessed darting levels 286 (Figure 3Di). There was a main effect of sex (F(1,34)=4.816, p=0.0351), but no effect of trial 287 (F(6.957, 236.6)=0.6941, p=0.6762) and no significant interaction (F(19, 646)=1.083, p=0.3640). 288 Post hoc Sidak's multiple comparisons test showed no significant differences between males 289 and females for any trial. For the extinction memory test one day later (Figure 3Dii), there was a 290 significant cue X sex interaction (F(3.102)=4.447, p=0.0056), as well as a main effect of both 291 cue (F(2.013, 68.44)=4.248, p=0.018) and sex (F1, 34)=4.834, p=0.0348). Females showed a 292 significantly higher dart rate than males during the fear cue (post hoc Sidak's multiple 293 comparisons test, p<0.05), which was also significantly higher than the dart rate to the reward 294 and safety cues in the females (post hoc Sidak's multiple comparisons test, p<0.05). However, 295 even though statistically significant, the amount of darting during the fear cue in females was 296 very low (0.05-0.4 in extinction training and 0.065 in the extinction memory test), and therefore 297 no clear conclusions can be made regarding darting and extinction in this study.

298

299 3.4 Shock reactivity in males versus females

300 To exclude possible sex differences in pain sensitivity and footshock perception, a separate 301 cohort of 8 male and 8 age-matched female rats received 11 unsignaled footshocks of 302 increasing intensities (0.3 mA, 0.35 mA, 0.4 mA, 0.45 mA, 0.5 mA, 0.55 mA, 0.6 mA, 0.7 mA, 303 0.8 mA, 0.9, 1.0 mA) with an inter-stimulus interval of 2 min. Freezing increased as a function of 304 shock intensities (Figure 4A; 2-way RM ANOVA; F(11,121)=25.9, p<0.0001). No main effects of 305 sex (F(1,121)=0.2871, p=0.6027) or sex by shock (F(11,121)=1.413, p=0.1754) were observed. 306 Our experiments utilized a shock intensity of 0.5mA throughout this study. For this particular 307 intensity, we also noted the number of rats that jumped or darted in response to a 0.5mA shock 308 (Figure 4B.C). No sex differences in the number of rats jumping in response to the 0.5mA 309 footshock were observed (χ^2 : p>0.9). The number of female rats darting after the 0.5mA 310 footshock was higher than males, but not significantly (χ^2 : p =0.0769), with five of the eight 311 female rats tested exhibiting the behavior. A higher number of females darting in response to 312 the footshock in this test would still not explain the lack of conditioned inhibition of freezing in 313 the females, as freezing levels at 0.5mA was slightly lower than the males (Figure 4A). Our

314 results do not definitively show, but do suggest, that females may be more likely to respond to a 315 footshock with a darting response.

316

317 4. Discussion

318 In this study, we show females exhibit a significantly different behavioral profile than males in a 319 task that tests for reward, fear and safety cue discrimination, as well as conditioned inhibition 320 and extinction. Female Long Evans rats showed more reward seeking early in training and 321 persistently high freezing levels to the fear cue when in the presence of a safety cue or after 322 fear extinction. Darting behavior in the females late in training showed conditioned inhibition of 323 this behavior in the presence of a safety cue, suggesting the females are able to discriminate 324 between the fear and safety cues but do not suppress their freezing response. This data adds to 325 the growing body of evidence of sex differences in fear regulation and highlights the advantages 326 of using more complex learning paradigms with additional behavioral measurements.

327

328 Even though studies including female subjects have been proportionally low, several studies 329 have reported clear sex differences in fear regulation. Most of these are consistent with our 330 findings of reduced discrimination between fear and safety signals. For instance, female mice 331 show more generalization of fear to novel and safe contexts compared to males, and with this 332 generalization there is a concurrent increase in basal amygdala activity (Keiser et al., 2017). 333 Male and female rats also respond differently to the controllability of a stressor. Males display 334 reduced fear during escapable stress versus inescapable stress whereas females exhibit no 335 beneficial effects of perceiving a stressor as escapable and controllable (Baratta et al., 2018). 336 The buffering effects seen in these males were linked to prelimbic cortical neurons projecting to 337 the dorsal raphe nucleus, which do not appear to be engaged in females. Females displaying a 338 similar fear response to both inescapable and escapable stress is similar to our findings of 339 females showing equivalent freezing levels to the fear cue in the presence or absence of a 340 safety cue, in that there were no buffering effects seen by the safety cue. It appears that 341 females do not downregulate their fear response in situations cued as safe. 342 343 Our data showing an increase in darting behavior in female rats as the number of fear cue-344 footshock trials increase is consistent with another report using female rats in a fear conditioning 345 and extinction paradigm (Gruene et al., 2015). Like us, Gruene et al (2015) also show darting

- 346 levels increase as learning about the fear cue advances. Compared to us, Gruene et al (2015)
- 347 report notably higher darting frequencies, which is most likely due to the differences in shock

intensities and number of trials; our study used 4 trials of 0.5mA per day for 4 days compared to their study using 7 trials of 0.7mA on one day. Our study also includes reinforced reward trials within the same sessions as the fear cue-footshock trials, which could alter the contextual expectations of the training session and reduce overall darting levels. It would be interesting in future studies to identify what leads a female to become a 'darter' versus 'non-darter'. As darting is a more active response compared to freezing, the circuits engaged during potential threats would likely be different in these two populations.

355

356 Our findings showing a lack of conditioned inhibition of freezing in females appear to be 357 inconsistent with a recent study demonstrating a lack of sex differences in conditioned inhibition 358 of freezing (Foilb et al., 2018). This is likely due to differences in our respective protocols. First, 359 their footshock intensity was 1.2mA, resulting in freezing levels >90% during the fear cue. As 360 footshock intensity and number of trials are consistently inconsistent across studies, it would be 361 interesting to assess if freezing and darting levels in females follow a linear trend with increasing 362 training intensity, or if there is instead a possibly U-shaped relationship. Foilb et al (2018) also 363 used separate presentations of the fear cue and safety cue throughout training and employed 364 the fear+safety cue summation test during recall, whereas we include fear+safety trials as part 365 of the training. In contrast, another study has shown females discriminate equally to males early 366 in training but then generalize their fear response to the safety cue with continued training (Day 367 et al., 2016). While the females in our study clearly showed equivalent freezing levels to both 368 the fear and fear+safety cues at all time points throughout training, they did not increase their 369 freezing levels to the safety cue when presented alone. And, lastly, our paradigm, unlike others, 370 includes reinforced reward trials during the training of fear and safety cues, which would change 371 the context from a 'threat-no threat' situation to a 'threat-no threat-reward' situation, inducing 372 approach behaviors on top of defensive behaviors.

373

374 Altogether, the data paints a consistent picture of females showing heightened fear responses 375 to cues signaling safety, mimicking the clinical picture in women (Gamwell et al., 2015; Lonsdorf 376 et al., 2015). The presentation of a safety signal not only decreases fear, but also stimulates 377 opposing neuronal activity. Field potential recordings in the striatum during safety signal 378 presentation has shown that brain regions dealing with approach and reward become activated 379 (Rogan et al., 2005). These findings have also been translated to using safety signals to 380 overcome anhedonia in rats (Pollak et al., 2008), showing that safety signals may also be 381 regulating emotion in addition to conditioned behavior (Foilb & Christianson, 2018).

382

383 In our study, females consistently showed elevated reward seeking behavior during the reward 384 cue compared to males beginning in the second reward pre-training session. This data appears 385 consistent with reward studies showing significant sex differences in response to sucrose, with 386 females willing to work more for sucrose in a progressive ratio paradigm (Tapia et al., 2019), 387 and in response to drugs of abuse, with female rats consistently self-administering drugs more 388 rapidly than males (Becker & Koob, 2016). The increased reward seeking in females seen in our 389 study remained until the end of the first DC session at which point they were equivalent to the 390 males. Interestingly, DC1 is the first time the animals are exposed to footshock. Taking into 391 account the lack of conditioned inhibition of freezing in the females, the females may no longer 392 be as motivated to seek rewards in the face of adverse footshocks. This would be consistent 393 with the report that female rats sacrifice their metabolic needs in order to avoid shocks more 394 than males (Pellmanet al., 2017).

395

396 Numerous sex differences have been reported in the functioning of the stress neuropeptide, 397 corticotropin-releasing factor (CRF), with differences in receptor expression, distribution, 398 trafficking and signaling (reviewed in (Bangasser & Wiersielis, 2018)). The majority of these 399 differences lead to enhanced CRF efficacy in females which may lead to heightened sensitivity 400 to stressors in females. Recently, the gene for CRH receptor 1 (CRHR1) has been identified as 401 a possible candidate gene for mood and anxiety disorders. Weber et al. (2016) have shown that 402 carrying the CRHR1 minor rs17689918 allele increases the risk for panic disorders in women. 403 Patients carrying this risk allele also demonstrate more generalization of fear to a safety cue, 404 increased amygdala activation during the safety cue and decreased frontal cortex activation with 405 discriminative fear conditioning. Thus, aberrant CRF signaling can lead to sustained fear under 406 conditions cued as safe and can be manifested by changes in neural activity in the amygdala 407 and frontal cortex.

408

Neural activity in the amygdala and prefrontal cortex has been shown by our lab to also play a critical role in effective discriminative conditioning in male rats. We have previously identified neurons in the basolateral amygdala (BLA) that discriminate among safety, fear and reward cues in male rats (Sangha et al., 2013); our future experiments will test if females show the same discriminative neurons. Using reversible pharmacological inactivations in male rats, we have also demonstrated that the infralimbic prefrontal cortex (IL) is necessary for suppression of conditioned fear during a safety cue and the prelimbic prefrontal cortex (PL) is necessary for

416 fear expression and discriminatory reward seeking (Sangha, Robinson, et al., 2014). These

- 417 results indicate that activating the IL in the females may improve conditioned inhibition to the
- 418 combined fear and safety cues. Our results with male rats also show that manipulating D1-
- 419 receptor mediated dopamine activity in the BLA disrupts suppression of conditioned fear (Ng et
- 420 al., 2018), implicating dopaminergic ventral tegmental area (VTA) neurons projecting to the BLA
- 421 in safety-fear-reward discrimination.
- 422
- 423 Our findings are consistent with human studies where females show less discrimination
- 424 between the fear and safety signals than males (Gamwell et al., 2015; Lonsdorf et al., 2015),
- 425 which may reflect underlying mechanisms of increased prevalence for anxiety and stress-
- 426 related disorders in women. For example, a deficiency in effective safety signal processing has
- 427 been linked to Post-traumatic Stress Disorder (Jovanovic et al., 2009, 2010), panic disorder
- 428 (Gorka et al., 2014), and anxiety (Lissek et al., 2005), all disorders with a higher incidence in
- 429 women than men (Mclean et al., 2011). In our paradigm, females show a significantly different
- 430 behavioral profile than males that is consistent with the clinical picture, thus making it a great
- 431 tool to test for the neurobiological mechanisms underlying these sex differences.
- 432

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562 **7. Figure Legends**

563 Figure 1. Females show increased reward seeking in response to the reward cue. A)

564 Schematic depicting experimental outline. During reward pre-training, rats (16 males, 20 565 females) received 25 cue-sucrose pairings across 5 separate sessions. **B)** Averaged percent 566 time spent in the reward port during the five reward pre-training sessions (R1-5). Females spent 567 significantly more time in the port compared to males during R2, R3 and R5. **C)** Averaged 568 latency to enter the port after cue onset (in seconds). Females entered the port significantly 569 sooner than males during R3-5. Means +/- SEM. *p<0.05, **p<0.01.

570

571 Figure 2. Females do not show inhibition of conditioned freezing in the presence of the

572 safety cue. A) Schematic depicting experimental outline. During the 4 DC sessions, rats (16 573 males. 20 females) were presented with four types of cued trials; reward cue-sucrose, fear cue-574 shock, fear+safety cue with no footshock and the safety cue presented alone without footshock. 575 B) Averaged percent time spent in the port during each cue across the 4 DC sessions. Both 576 males and females showed significantly higher reward seeking during the reward cue compared 577 to all other cues during every DC session. During DC1, females showed significantly higher 578 reward seeking to the reward cue compared to males. C) Averaged percent time spent freezing 579 during each cue across the 4 DC sessions. During DC3 and DC4, males showed significantly 580 lower freezing to the fear+safety cue (and reward and safety cues) when compared to the fear 581 cue. Females did not show significant inhibition of conditioned freezing to the fear+safety cue 582 compared to the fear cue during any DC session. Females also showed significantly higher 583 freezing to the fear+safety cue compared to males during every session. D) Darting behavior 584 during each cue across the 4 DC sessions. During DC4 females showed significantly more darts 585 than males during the fear and fear+safety cues. Means +/- SEM. # p<0.05, ####p<0.0001 586 within sex, between cue comparison; * p<0.05, **p<0.01, ***p<0.001, ****p<0.0001 within cue,

587 between sex comparison.

588

Figure 3. Females do not show significant extinction of fear. A) Schematic depicting experimental outline. During extinction training both the reward and fear cues are presented in the same session without sucrose or footshock. During the test for extinction memory 1 day later all cues are presented without sucrose or footshock. Bi) Averaged percent time spent in the port during each reward cue presentation during extinction training. No significant differences were found between males and females during extinction training. Bii) Averaged percent time spent in the port during each cue 1 day after extinction training. Females spent 596 significantly more time in the port than males during the safety cue. Ci) Averaged percent time 597 spent freezing during each fear cue presentation during extinction training. Compared to the first 598 trial of extinction, males showed significantly reduced freezing during trials 8, 9 and 11-20. 599 Freezing levels for females did not significantly decrease at any point in extinction training, with 600 the exception of trial 19. #p<0.05, compared to trial 1. Cii) Averaged percent time spent freezing 601 during each cue 1 day after extinction training. Males showed evidence of fear cue extinction 602 retention. Females froze significantly more than males during the fear and fear+safety cues. Di) 603 Averaged darting during each fear cue presentation during extinction training. No significant 604 post hoc differences found between males and females during extinction training. Dii) Averaged 605 darting during each cue 1 day after extinction training. Females had a significantly higher dart 606 rate than males during the fear cue, which was also significantly higher than the reward and 607 safety cues in females. Means +/- SEM. #p<0.05, ####p<0.0001 within sex, between cue/trial 608 comparisons. *p<0.05, **p<001, ****p<0.0001 within cue, between sex comparisons. 609 610 Figure 4. No significant differences in shock reactivity between age-matched male and 611 female rats. A) Male and female rats (n=8 each) were subjected to increasing footshock 612 intensities from 0.3mA to 1.0mA. No significant differences in freezing levels (means +/- SEM)

613 were detected between males and females after each shock presentation. The box around the

614 data at 0.5mA indicates the intensity used for the experiments in this study. There were no

615 significant differences in the number of males or females who jumped (B) or darted (C) in

616 response to the 0.5mA shock.

Reward seeking				
Session	Cue x Sex effects	Main effect of cue	Main effect of sex	
DC1	F(3,102) = 3.472,	F(3,102) = 95.16,	F(1,34) = 9.827,	
	p=0.0189	p<0.0001	p=0.0035	
DC2	F(3,102) = 0.7742,	F(3,102) = 227.9,	F(1,34) = 4.69,	
	p=0.5110	p<0.0001	p=0.0374	
DC3*	F(3,90) = 0.6512,	F(3,90) = 117,	F(1,30)=1.041,	
	p=0.5843	p<0.0001	p=0.3157	
DC4	F(3,102) = 2.255,	F(3,102) = 181.2,	F(1,34) = 2.453,	
	p=0.0864	p<0.0001	p=0.1266	
Freezing				
Session	Cue x Sex effects	Main effect of cue	Main effect of sex	
DC1	F(3,102) = 2.245,	F(3,102) = 31.82,	F(1,34) = 5.045,	
	p=0.0876	p<0.0001	p=0.0313	
DC2	F(3,102) = 4.075,	F(3,102) = 103.4,	F(1,34) = 6.621,	
	p=0.0089	p<0.0001	p=0.0146	
DC3*	F(3,90) = 2.9,	F(3,90) = 151.3,	F(1,30)=9.719,	
	p=0.0393	p<0.0001	p=0.0040	
DC4	F(3,102) = 4.889,	F(3,102) = 198.9,	F(1,34) = 8.294,	
	p=0.0032	p<0.0001	p=0.0068	
Darting				
Session	Cue x Sex effects	Main effect of cue	Main effect of sex	
DC1	F(3,102) = 1.98,	F(2.022,68.73) =	F(1,34) = 4.146,	
	p=0.1216	2.388,	p=0.0496	
		p=0.0989		
DC2	F(3,102) = 1.134,	F(1.923, 65.38) =	F(1,34) = 3.667,	
	p=0.3390	9.377,	p=0.0640	
		p=0.0003		
DC3	F(3,102) = 0.9158	F(1.586,53.93) =	F(1,34)=0.9579,	
	p=0.4361	18.96,	p=0.3346	
		p<0.0001		
DC4	F(3,102) = 10.65,	F(1.684, 57.26) =	F(1,34) = 13.34,	
	p<0.0001	15.65,	p=0.0009	
		p<0.0001		

*video files for 4 females were corrupted for this session (n=16 females, 16 males)

Table 1. Summary of two-way repeated-measures ANOVA analyses for reward seeking, freezing and darting behaviors during the four discriminative conditioning (DC) sessions.







