1 Title: Short-term social isolation acts on hypothalamic neurons to promote social

2 behavior in a sex- and context-dependent manner.

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7 Abstract

8 Social animals, including both humans and mice, are highly motivated to engage in 9 social interactions. Short-term social isolation increases social motivation and promotes 10 social behavior, but the neural circuits through which it does so remain incompletely 11 understood. Here, we sought to identify neurons that promote social behavior in single-12 housed female mice, which exhibit increased rates of social investigation, social ultrasonic 13 vocalizations (USVs), and mounting during same-sex interactions that follow a period of 14 short-term (3-day) isolation. We first used immunostaining for the immediate early gene Fos 15 to identify a population of neurons in the preoptic hypothalamus (POA) that increase their 16 activity in single-housed females following same-sex interactions (POA_{iso} neurons). TRAP2-17 mediated chemogenetic silencing of POA_{iso} neurons in single-housed females significantly 18 attenuates the effects of short-term isolation on social investigation and USV production and 19 also tends to reduce mounting. In contrast, caspase-mediated ablation of POA_{iso} neurons in 20 single-housed females robustly attenuates mounting but has no effect on social investigation 21 or USV production. Optogenetic activation of POA_{iso} neurons in group-housed females 22 promotes USV production but does not recapitulate the effects of short-term isolation on 23 social investigation and mounting. To understand whether a similar population of POA_{iso} 24 neurons promotes social behavior in single-housed males, we performed Fos 25 immunostaining in single-housed males following either same-sex or opposite-sex social 26 interactions. These experiments revealed a population of POA neurons that increase Fos 27 expression in single-housed males following opposite-sex, but not same-sex, interactions. 28 Chemogenetic silencing of POA_{iso} neurons in single-housed males during interactions with 29 females tends to reduce mounting but does not decrease social investigation or USV 30 production. These experiments identify a population of hypothalamic neurons that promote 31 social behavior following short-term isolation in a sex- and social context-dependent manner. 32 Keywords: social isolation, ultrasonic vocalization, preoptic hypothalamus, mounting,

33 female

34 Introduction

35 Humans and other social mammals find social interactions rewarding and are highly 36 motivated to seek out social connections. Consequently, the experience of social isolation is 37 aversive and impacts both our brains and our behaviors. While long-term isolation can lead 38 to the emergence of anti-social behaviors in both humans and rodents (An et al., 2017; 39 Arrigo and Bullock, 2008; Check et al., 1985; Hossain et al., 2020; Killgore et al., 2021; Ma 40 et al., 2011, 2022; Machimbarrena et al., 2019; Matsumoto et al., 2005; Mears and Bales, 41 2009; Reid et al., 2022; Toth et al., 2011; Valzelli, 1973; Weiss et al., 2004; Wiberg and 42 Grice, 1963; Zelikowsky et al., 2018), short-term isolation typically increases levels of social 43 motivation and promotes social-seeking behaviors (Baumeister and Leary, 1995; Cacioppo 44 et al., 2006; Cacioppo and Cacioppo, 2018; House et al., 1988; Lee et al., 2021; Niesink and 45 van Ree, 1982; Panksepp and Beatty, 1980; Zhao et al., 2021). Alterations in social 46 motivation are characteristic of many neurodevelopmental disorders, including autism 47 spectrum disorder (Chevallier et al., 2012; Clements et al., 2018). How short-term social 48 isolation acts on the brain to promote social behavior remains incompletely understood.

49 Mesolimbic circuits play an important role in regulating social motivation and social 50 reward, during courtship as well as during same-sex interactions (Bariselli et al., 2018; Dai et 51 al., 2022; Dölen et al., 2013; Gunaydin et al., 2014; Hung et al., 2017; Love, 2014; Melis et 52 al., 2022; Resendez et al., 2020; Robinson et al., 2011; Solié et al., 2022; Tang et al., 2014; 53 Walum and Young, 2018; Xiao et al., 2017). In line with their role in regulating social 54 behavior, changes in the function of mesolimbic circuits have been reported following long-55 term (weeks-long) social isolation and/or early-life social isolation (McWain et al., 2022; 56 Musardo et al., 2022; Tan et al., 2021; Yorgason et al., 2016). The neural circuit changes 57 that mediate the effects of short-term isolation on social motivation in adult animals are 58 comparatively less explored, but here as well, recent studies in both humans and rodents 59 have implicated changes in various populations of midbrain dopamine neurons (Inagaki et 60 al., 2016; Matthews et al., 2016; Tomova et al., 2020). Beyond its effects on mesolimbic 61 circuits, whether social isolation acts on additional neuronal populations to promote social 62 interaction is unknown.

In recent work, we found that short-term (3-day) social isolation exerts robust effects
on the social behaviors of C57BL/6J female mice (Zhao et al., 2021). Relative to grouphoused females, single-housed females that subsequently engaged in same-sex interactions
exhibited increased rates of social investigation, increased rates of USVs, and were also
observed to mount female social partners, a behavior never observed in pairs of group-

68 housed females (Zhao et al., 2021). The robust effect of short-term isolation on these three 69 aspects of female social behavior provides a powerful paradigm to identify neurobiological 70 changes that mediate the effects of short-term isolation on social behavior. In the current 71 study, we combined this behavioral paradigm with Fos immunostaining and the TRAP2 72 activity-dependent labeling approach (Allen et al., 2017; DeNardo et al., 2019) to identify and 73 characterize a population of neurons in the preoptic hypothalamus that increase their activity 74 in single-housed females following same-sex social interactions (i.e., POA_{iso} neurons). We 75 next asked whether silencing or ablation of POA_{iso} neurons attenuates the effects of short-76 term isolation on female social behavior, and whether artificial activation of POA_{iso} neurons 77 in group-housed females mimics the effects of short-term isolation on female social 78 behavior. Finally, we extended a subset of these experiments to single-housed males 79 engaged in opposite-sex and same-sex interactions, to understand whether short-term 80 isolation acts on the POA to promote social behavior in a manner that depends on either sex 81 or social context. This study identifies novel neurobiological mechanisms through which 82 short-term social isolation acts on the brain to promote social interaction. Our findings also 83 add to an emerging literature indicating that the POA regulates not only sexual behavior but 84 also female social behavior during same-sex interactions.

85 Results

Neurons in the preoptic hypothalamus increase their activity in socially isolated female mice following same-sex social interactions

88 To identify changes in neuronal activity that may underlie the effects of short-term 89 isolation on female social behavior, we performed immunostaining for the immediate early 90 gene Fos in brain sections collected from group-housed and single-housed (3-days) subject 91 females following 30-minute social encounters in their home cages with a novel, group-92 housed visitor female (Fig. 1A). In line with our previous behavioral findings (Zhao et al., 93 2021), we observed that single-housed female residents spent more time investigating 94 visitors (Fig. 1B; Mann-Whitney U test, p = 0.001) and in many trials mounted visitors, a 95 behavior that was not observed in group-housed residents (Fig. 1C; 0 of 12 group-housed 96 residents and 10 of 13 single-housed residents mounted visitors; Mann-Whitney U test for 97 difference in total mounting duration, p < 0.001; see Table S1 for complete statistical 98 details). Female pairs that contained a single-housed resident also produced higher rates of 99 ultrasonic vocalizations (USVs) than pairs with a group-housed resident (Fig. 1D; Mann-100 Whitney U test, p < 0.001). Although either female in a dyad can produce USVs (Warren et 101 al., 2020), the robust effects of short-term isolation on the non-vocal social behaviors of 102 single-housed females suggest that at least some of the elevation in USV rates is driven by

increased USV production by the single-housed resident. Given the robust effects of short-

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104 term isolation on these three aspects of female social behavior, we focused our analyses on 105 two hypothalamic regions implicated in regulating these behaviors: the preoptic area (POA), 106 which regulates social approach (McHenry et al., 2017), social reward (Hu et al., 2021), 107 mounting (Floody, 1989; Karigo et al., 2021; Wei et al., 2018) and USV production (Chen et 108 al., 2021; Gao et al., 2019; Green et al., 2018; Karigo et al., 2021; Michael et al., 2020); and 109 the ventromedial hypothalamus (VMH), which regulates mounting (Hashikawa et al., 2017; 110 Karigo et al., 2021; Lee et al., 2014; Liu et al., 2022). We also examined Fos expression 111 within the caudolateral periaqueductal gray (PAG), based on the well-established role of this 112 region in the control of vocalization in vertebrates and USV production in mice (Chen et al., 113 2021; Jürgens, 1994; Michael et al., 2020; Tschida et al., 2019). To test whether any 114 observed differences in Fos expression in these three regions were associated with 115 isolation-induced changes in social behavior rather than baseline differences between 116 groups, we also measured Fos expression in the POA, the VMH, and the PAG of group-117 housed and single-housed females that did not engage in social interaction with novel 118 female visitors (Fig. 1E-F; Fig. S1). 119 These analyses revealed that baseline levels of Fos expression within the POA and 120 the VMH did not differ between group-housed and single-housed females (Fig. 1F, left and 121 middle, open bars; two-way ANOVA to analyze Fos expression within each brain region, 122 factor 1 = housing status, factor 2 = social interaction, followed by post-hoc Tukey's HSD 123 tests). Following social interactions with novel female visitors, single-housed females 124 exhibited robust increases in Fos expression within the POA (Fig. 1F, left; p < 0.001) but not 125 within the VMH (Fig. 1F, middle; p > 0.05). In contrast, Fos expression within these two brain 126 areas did not increase significantly in group-housed females that interacted with novel 127 female visitors (Fig. 1F; p > 0.05 for both comparisons). Similar to the POA, baseline Fos 128 expression within the PAG did not differ between group-housed and single-housed females 129 (p > 0.05), and only single-housed females displayed increased PAG Fos expression 130 following social interactions with novel female visitors (Fig. 1F, right; p < 0.001), a finding 131 that further supports the idea that single-housed females increase USV production during 132 same-sex interactions. POA Fos expression was significantly and positively correlated with 133 the total amount of time spent in resident-initiated investigation for both group-housed and 134 single-housed females (Fig. S1C, left; linear regression, p < 0.05), as well as with the total 135 number of mounting bouts produced by single-housed resident females (Fig. S1C, middle; p 136 < 0.01). In both group-housed and single-housed female residents, Fos expression tended 137 to correlate positively with total USVs, but these relationships were not significant (Fig. S1C, 138 right; p > 0.05 for both comparisons; see Fig. S1D-E for relationships of VMH Fos and PAG

139 Fos to vocal and non-vocal social behaviors). In summary, POA Fos expression increases

selectively in single-housed females following social interactions, and levels of POA Fos
expression are also well related to the production of specific types of social behaviors by
single-housed females.

143 To ask whether the effects of short-term isolation on female social behavior and POA 144 Fos expression are long-lasting, we measured social behaviors of female residents at three 145 timepoints: (1) on day 0, when female subjects were still group-housed; (2) on day 3, after 146 female subjects had been single-housed for 3 days; and (3) on day 17, after half of the 147 subject females had been re-group-housed with their same-sex siblings for two weeks and 148 the other half of the subject females remained single-housed for two weeks (Figs. 1G-I). 149 Brains of re-group-housed and 14-day single-housed subject females were collected after 150 the day 17 social interaction, and Fos expression within the POA was examined (Fig. 1K). 151 Consistent with our earlier findings, rates of social investigation and USV production 152 significantly increased following 3 days of social isolation (Fig. 1G, I; one-way ANOVAs with 153 repeated measures; p < 0.05 for day 0 vs. day 3 in both groups for both behaviors). 154 Following re-group-housing, time spent in social investigation and rates of USV production 155 tended to decrease to pre-isolation levels (Fig. 1G, I, top plots; p = 0.06 for day 0 vs. day 17 156 investigation time and day 0 vs. day 17 total USVs in re-group-housed females). In contrast, 157 females that were single-housed for 14 days continued to spend increased time in social 158 investigation (Fig. 1G, bottom plot; p < 0.05 for day 0 vs. day 3 investigation and for day 0 159 vs. day 17 investigation), and pairs containing 14-day single-housed residents continued to 160 produce elevated rates of USVs (Fig. 1I, bottom plot; p < 0.05 for day 0 vs. day 3 USVs and 161 for day 0 vs. day 17 USVs). Time spent mounting tended to follow the same trends as rates 162 of social investigation and USV production in re-group-housed and 14-day single-housed 163 females (Fig. 1H). Along with the attenuation of female social behaviors following re-group-164 housing, we also found that POA Fos expression was significantly lower in re-group-housed 165 females relative to 14-day single-housed females (Fig. 1K; t-test, p < 0.001). These findings 166 support the idea that changes in female social behavior following short-term isolation are 167 reversible and are accompanied by decreased POA Fos expression. Hereinafter, we refer to 168 the population of POA neurons that increase Fos expression in single-housed females that 169 have engaged in same-sex interactions as POA_{iso} neurons, and we next conducted 170 experiments to test whether functional manipulations of POA_{iso} neuronal activity impact the 171 effects of short-term isolation on female social behavior.

172 Chemogenetic inhibition of POA_{iso} neurons attenuates the effects of social isolation 173 on female social investigation and USV production

If increased activity of POA_{iso} neurons contributes to the effects of short-term
 isolation on female social behavior, one prediction is that reducing the activity of POA_{iso}

176 neurons in single-housed females will attenuate the effects of isolation on female social 177 behavior. To test this idea, we employed the TRAP2 activity-dependent labeling strategy to 178 chemogenetically silence POA_{iso} neurons in single-housed females during social interactions 179 with novel, group-housed female visitors (Fig. 2A). Briefly, the POA of TRAP2 female mice 180 was injected bilaterally with a virus driving the Cre-dependent expression of the inhibitory 181 DREADDs receptor hM4Di. Three weeks later, females were single-housed for 3 days and 182 then given a 30-minute social encounter with a novel, group-housed female visitor in their 183 home cage. Following the social interaction, resident females were given an I.P. injection of 184 4-hydroxytamoxifen (4-OHT), which drives the transient expression of Cre recombinase in 185 recently active neurons and thereby enables the expression of hM4Di in POA_{iso} neurons. 186 Subject females remained single-housed for an additional 24 hours and then were re-group-187 housed with siblings for 2 weeks. Subject females were then single-housed a second time 188 for 3 days and subsequently given a 30-minute same-sex interaction following I.P. injection 189 of either saline (control) or clozapine-n-oxide (CNO) (saline and CNO tests were run 3 days 190 apart, and the order was counterbalanced across experiments).

191 Comparison of the social behaviors of single-housed females between CNO and 192 saline sessions revealed that chemogenetic silencing of POA_{iso} neurons significantly 193 reduced resident-initiated investigation (Fig. 2B; N = 12; red points; two-way ANOVA with 194 repeated measures on one factor; p < 0.01). Inhibition of POA_{iso} neurons also tended to 195 reduce mounting, although this effect was not statistically significant (Fig. 2D; Kruskal Wallis 196 test performed on difference in mounting time (CNO-saline) for each group; p > 0.05). 197 Finally, inhibition of POA_{iso} neurons significantly reduced USV production (Fig. 2D; two-way 198 ANOVA with repeated measures on one factor; p < 0.01). In contrast, CNO treatment did not 199 affect the production of any of these social behaviors in single-housed females with GFP 200 expressed in POA_{iso} neurons (Fig. 2B-D; N = 14; black points; p > 0.05 for all CNO vs. saline 201 comparisons in the POA_{iso} FLEX-GFP control group). To investigate the specificity of these 202 effects to chemogenetic silencing of POA_{iso} neurons, we also performed control experiments 203 in which activity-dependent chemogenetic silencing was performed caudal to the POA within 204 the anterior hypothalamus (AH) (Figs. 2B-D; N = 12; brown points) or within the VMH (Fig. 205 2B-D; N = 5; gray points). No significant effects of CNO treatment on resident-initiated 206 investigation, mounting, or USV rate were observed in these control groups (Figs. 2C-D; p > 207 0.05 for all). The effect of chemogenetic inhibition of POA_{iso} neurons to decrease female 208 social behavior also cannot be attributed to an overall decrease in movement (Fig. 2E; p > 209 0.5 for difference in movement between saline and CNO sessions; see Methods). In 210 summary, we demonstrate that chemogenetic silencing of POA_{iso} neurons attenuates 211 isolation-induced changes in social behavior in female mice.

Ablation of POA_{iso} neurons attenuates the effects of social isolation on female mounting

214 In previous work investigating the role of the POA in regulating rodent social 215 behaviors, studies have reported different effects on behaviors according to whether they 216 employed reversible or irreversible neuronal silencing strategies. Studies that used 217 chemogenetic or optogenetic methods to reversibly silence genetically-defined subsets of 218 POA neurons report decreases in both USV production in males (Chen et al., 2021; Karigo 219 et al., 2021) and in mounting in males and females during interactions with female social 220 partners (Gao et al., 2019; Karigo et al., 2021). In contrast, studies employing caspase-221 mediated ablation of genetically-defined subsets of POA neurons (Gao et al., 2019; Wei et 222 al., 2018) or electrolytic lesions of the POA (Bean et al., 1981) report decreased mounting 223 but no effects on rates of USV production. To test whether permanent ablation of POA_{iso} 224 neurons attenuates the effects of social isolation on female behavior in a manner similar to 225 the effects of chemogenetic inhibition, we used the TRAP2 activity-dependent labeling 226 strategy to express caspase in and to thereby ablate POA_{iso} neurons (Fig. 3A; see Methods). 227 Vocal and non-vocal social behaviors of resident females were compared pre- and post-228 ablation, and the same measurements were made in control females expressing GFP in 229 POA_{iso} neurons.

230 In contrast to the effects of chemogenetic inhibition of POA_{iso} neurons, we found that 231 caspase-mediated ablation of POA_{iso} neurons did not affect rates of social investigation in 232 single-housed females, although both experimental and control females spent more time 233 investigating visitors in the post-4-OHT session (Fig. 3B; two-way ANOVA with repeated 234 measures on one factor; p > 0.05 for main effect of group, p < 0.01 for main effect of time, p 235 > 0.05 for interaction effect). Ablation of POA_{iso} neurons also failed to reduce USV 236 production in pairs containing single-housed females (Fig. 3D; two-way ANOVA with 237 repeated measures on one factor; p > 0.05 for pre-4-OHT vs. post-4-OHT USV rates in 238 POA_{iso}-caspase females). Notably, ablation of POA_{iso} neurons significantly reduced 239 mounting in single-housed females (Fig. 3C; Mann Whitney U test performed on the 240 difference in mounting time (post-4-OHT - pre-4-OHT, p = 0.01). Taken together with our 241 chemogenetic inhibition data, these results show that both reversible inhibition or irreversible 242 ablation of POA_{iso} neurons in single-housed female mice attenuates the effects of short-term 243 isolation on mounting behavior, whereas only chemogenetic inhibition of POA_{iso} neurons 244 attenuates the effects of short-term isolation on female social investigation and USV 245 production.

246 **Optogenetic activation of POA**_{iso} neurons elicits USV production

247 To understand whether artificial activation of POA_{iso} neurons can recapitulate the 248 effects of short-term isolation on female social behavior, we assessed the effects of 249 optogenetic activation of POA_{iso} neurons on the social behaviors of group-housed females. 250 The TRAP2 strategy was used to express either channelrhodopsin (ChR2) or GFP in POA_{iso} 251 neurons (Fig. 4A; see Methods), and females were re-group-housed for two weeks before 252 beginning optogenetic activation experiments. The effects of optogenetically activating 253 POA_{iso} neurons were first assessed for each subject female in a 5-minute solo session, in 254 which the female was tested alone in a behavior chamber while pulses of blue light were 255 delivered unilaterally to the POA (473 nm, 10 mW, 20-50 Hz, 10-20 ms pulses, 5-10s train 256 durations). The effects of optogenetically activating POA_{iso} neurons were then assessed for 257 each subject female in a 20-minute social session, in which a novel, group-housed female 258 visitor was added to the behavior chamber. The pair was allowed to interact in the absence 259 of optogenetic stimulation for the first and last 5 minutes of the social session, and pulses of 260 blue light were delivered to the POA of the subject female throughout the middle 10 minutes 261 of the session (Fig. 4A).

262 When POA_{iso}-ChR2 females were tested alone, we found that optogenetic activation 263 of POA_{iso} neurons elicited weak-to-moderate USV production in 4 of 8 females, but the 264 comparison of USV rates from pre-laser baseline to the laser stimulation period was not 265 significant at the level of the entire group (Fig. 4B; Mann Whitney U test performed on the 266 difference in USV rates (laser - pre-laser), p = 0.09). In POA_{iso}-GFP control females, laser 267 stimulation failed to elicit USV production (0 ± 0 USVs elicited in N = 6 POA_{iso-}GFP controls). 268 Interestingly, we found that when laser stimulation was applied during social sessions, 269 optogenetic activation of POA_{iso} neurons more readily elicited USV production than in solo 270 sessions (Fig. 4C; USVs elicited by blue laser stimulation in 7 of 8 POA_{iso-}ChR2 females; 271 Mann Whitney U test performed on the difference in USV rates (laser - pre-laser), p =272 0.006). Moreover, optogenetic activation elicited higher rates of USVs when applied at times 273 when subject females were in close proximity to visitor females (within 2 mouse body 274 lengths) as compared to times when the females were farther apart (mean increase in USV 275 rates from pre-laser to laser period was 2.96 ± 2.32 USVs/s for "near" stimulations, 1.84 ± 276 1.75 USVs/s for "far" stimulations; paired t-test performed on the difference in USV rates 277 (laser - pre-laser) for "far" vs. "near" stimulations; p = 0.02). In summary, optogenetic 278 activation of POA_{iso} neurons elicits USV production in group-housed females, and the 279 efficacy of this effect is modulated by social context and proximity to a social partner. 280 In contrast to the effects on USV production, optogenetic activation of POA_{iso} 281 neurons failed to extend the duration of social investigation bouts (paired t-test performed on 282 mean duration of social investigation bouts for each POA_{iso}-ChR2 female that overlapped

with laser stimulation vs. those that did not; p > 0.05). Moreover, optogenetic activation of POA_{iso} neurons only infrequently elicited mounting (activation elicited n =1 bout of mounting in N = 1 POA_{iso}.ChR2 female, n = 2 bouts of mounting in N = 1 POA_{iso}.ChR2 female, and n = 0 bouts of mounting in the remaining N = 7 POA_{iso}.ChR2 females). In summary, optogenetic activation of POA_{iso} neurons elicits USV production from group-housed females, particularly when female subjects are engaged in interactions with female visitors, but otherwise fails to recapitulate the effects of short-term isolation on the social behaviors of female mice.

290 Previous studies have found that USV production can be elicited in female and male 291 mice by artificial activation of VGAT⁺ POA neurons (Gao et al., 2019), Esr1⁺ POA neurons 292 (which are predominantly VGAT⁺) (Chen et al., 2021; Michael et al., 2020), as well as POA 293 neurons that send axonal projections to the caudolateral PAG (which are predominantly 294 VGAT⁺) (Chen et al., 2021; Michael et al., 2020). To ask to what extent POA_{iso} neurons 295 overlap with these previously described populations, we first evaluated the neurotransmitter 296 phenotype of POA_{iso} neurons by performing two-color in situ hybridization for c-fos mRNA 297 and vesicular GABA transporter (VGAT) mRNA and calculating the percentage of Fos⁺ POA 298 neurons that co-expressed VGAT. This analysis revealed that a majority of POA_{iso} neurons 299 are GABAergic (Fig. S2A-B; N = 4, 76 ± 8.8%). We next used the TRAP2 activity-dependent 300 labeling strategy to express GFP in POA_{iso} neurons and found GFP-positive axons within the 301 caudolateral PAG, indicating that at least some POA_{iso} neurons send axonal projections to 302 the PAG (Fig. S2C-D; see Methods). Finally, we combined retrograde tracing from the 303 caudolateral PAG with Fos immunostaining to quantify the percentage of PAG-projecting 304 POA neurons that increase Fos expression in single-housed females following same-sex 305 interactions. This experiment revealed that around 20% of PAG-projecting POA neurons 306 express Fos in single-housed females following same-sex interactions (Fig. S2E-F; N = 4307 females, percentage of tdTomato neurons that are Fos-positive = $18.3 \pm 2.9\%$). These 308 findings suggest that a subset of POA_{iso} neurons overlap with GABAergic, PAG-projecting 309 POA neurons that have been demonstrated in previous work to promote USVs via 310 disinhibition of excitatory PAG neurons important to USV production (Chen et al., 2021; 311 Michael et al., 2020).

POA neurons increase their activity in single-housed male mice following opposite sex but not same-sex social interactions

Given our findings that POA_{iso} neurons contribute to isolation-induced changes in the social behaviors of female mice, we next wondered whether a similar population of POA neurons contributes to isolation-induced changes in social behavior in male mice. To address these questions, we measured the vocal and non-vocal social behaviors of sexually naïve males, which were either group-housed with same-sex siblings or single-housed for 319 three days and then given a 30-minute social interaction with a novel, group-housed visitor. 320 To consider the effects of isolation on male social behavior in different social contexts, males 321 were given either a social encounter with a same-sex visitor (MM context) or with an 322 opposite-sex visitor (MF context). Following these social interaction tests, we collected the 323 brains of the subject males and performed immunostaining to measure Fos expression 324 within the POA. Consistent with our prior work (Zhao et al., 2021), we found that males 325 exhibit higher rates of social investigation, higher rates of mounting, and produce more 326 USVs during interactions with females than during same-sex interactions (Fig. 5A-C; p < 1327 0.05 for main effect of social context for all three behaviors). With respect to resident-328 initiated investigation, we found a significant main effect of housing, indicating that single-329 housed males spent more time investigating visitors during both opposite-sex and same-sex 330 interactions (Fig. 5A; two-way ANOVA, p = 0.02 for main effect of housing). In contrast, 331 single-housed males spent more time mounting female visitors than did group-housed 332 males, but there were no differences in mounting between single-housed and group-housed 333 males during same-sex interactions (Fig. 5B; two-way ANOVA, p < 0.05 for difference 334 between single-housed males interacting with females and all other groups). Similarly, there 335 was also a context-dependent effect of social isolation on male USV production, whereby 336 only single-housed males that interacted with female visitors exhibited increased USV 337 production relative to group-housed males (Fig. 5C; two-way ANOVA with post-hoc Tukey's 338 HSD tests; p < 0.001 for total USVs in single-housed MF vs. group-housed MF trials; p > 0.001339 0.05 for total USVs in single-housed MM vs. group-housed MM trials). The finding that short-340 term isolation exerts larger effects on male social behavior during subsequent opposite-sex 341 interactions relative to same-sex interactions is consistent with prior work (Zhao et al., 2021). 342 When we examined POA Fos expression in these four groups of males, we found that POA 343 Fos was significantly elevated in single-housed males following interactions with females 344 relative to the other three groups (Fig. 5D; two-way ANOVA, Tukey's post-hoc HSD tests; p 345 < 0.05 for difference in POA Fos between single-housed MF and all other groups). In 346 summary, the effects of short-term isolation on male social behavior are context-dependent, 347 and increased Fos expression within the POA is seen in single-housed males following 348 interactions with females, a context marked by increased male social investigation, 349 increased male mounting, and increased male USV production.

To test whether neural activity in male POA_{iso} neurons contributes to isolationinduced changes in male social behavior, we used the TRAP2 strategy to chemogenetically silence POA_{iso} neurons in single-housed males during social interactions with novel, grouphoused females (see Methods). The vocal and non-vocal behaviors of subject males were measured and compared during 30-minute social interactions following I.P. injection of either 355 saline or CNO. Control males were treated identically but were injected with a virus to drive 356 expression of GFP in POA_{iso} neurons. In contrast to our findings in females, chemogenetic 357 inhibition of male POA_{iso} neurons tended to reduce time spent mounting (Fig. 5G; Mann 358 Whitney U test performed on difference in mounting time (CNO - saline); p = 0.13 for 359 difference between groups) but did not change rates of resident-initiated social investigation 360 (Fig. 5F; two-way ANOVA with repeated measures on one factor; p = 0.01 for main effect of 361 group; p > 0.05 for main effect of drug and for interaction effect) and also did not affect USV 362 rates (Fig. 5H; two-way ANOVA with repeated measures on one factor; p > 0.05 for main 363 effects and interaction effect). In summary, we find that chemogenetic silencing of male 364 POA_{iso} neurons tends to reduce mounting during subsequent social interactions with females 365 but does not reduce social investigation or USV production, a pattern of results that differs 366 from the effects on single-housed female social behavior of chemogenetically silencing 367 female POA_{iso} neurons.

368 Discussion

369 In the current study, we identify and characterize a population of preoptic 370 hypothalamic neurons that contribute to the effects of short-term social isolation on the 371 social behaviors of mice. These POA_{iso} neurons exhibit increased Fos expression in single-372 housed female mice following same-sex social interactions, and this increase in Fos 373 expression scales positively with the time females spend investigating and mounting female 374 visitors and tends also to scale with rates of USVs. Chemogenetic silencing of POAiso 375 neurons attenuates the effects of social isolation on female social behavior, significantly 376 reducing social investigation and USV production while tending to reduce mounting. In 377 contrast, irreversible ablation of POA_{iso} neurons significantly reduces mounting in single-378 housed females but has no effect on rates of social investigation or USVs. Optogenetic 379 activation of POA_{iso} neurons partially recapitulates the effects of short-term isolation on 380 female behavior and promotes USV production in female mice, particularly during same-sex 381 social interactions and when in close proximity to female visitors. Finally, we extended our 382 analyses to male mice to understand whether similar POA neurons may mediate changes in 383 male social behavior following short-term isolation. We find that short-term isolation exerts 384 more robust effects on male behavior during subsequent interactions with females than 385 during subsequent interactions with males, and increased POA Fos expression is only seen 386 in single-housed males following social interactions with females. Interestingly, 387 chemogenetic silencing of these POA_{iso} neurons tends to reduce mounting but has no effect 388 on social investigation and USV production, in contrast to the effects of chemogenetically 389 silencing POA_{iso} neurons in females. Together, these experiments identify a population of

preoptic hypothalamic neurons that promote social behaviors in single-housed mice in amanner that depends on sex and social context.

392 An extensive body of past work has implicated the POA in the regulation of male 393 sexual behavior, including in the regulation of male courtship vocalizations in both rodents 394 and birds (Alger and Riters, 2006; Bean et al., 1981; Gao et al., 2019; Merari and Ginton, 395 1975; Riters, 2012; Riters et al., 2000; Riters and Ball, 1999; Wei et al., 2018). Past work 396 has also shown that activation of genetically-defined subsets of POA can elicit the 397 production of USVs in both male and female mice (Chen et al., 2021; Gao et al., 2019; 398 Karigo et al., 2021; Michael et al., 2020). However, whether the POA regulates natural USV 399 production in female mice remained unclear. In the current study, we demonstrate that 400 reversible silencing of POA neurons that increase their activity during same-sex interactions 401 decreases female USV production, indicating that the POA regulates the production of USVs 402 by single-housed females engaged in same-sex interactions. Whether these same neurons 403 regulate female USV production in other behavioral contexts, including in group-housed 404 females and in females interacting with male partners, remains an important open question. 405 In addition to attenuating USV production, we found that silencing of POA_{iso} neurons in 406 single-housed females significantly reduced social investigation of a same-sex partner and 407 also tended to reduce mounting. Our retrograde and anterograde tracing experiments 408 demonstrate that at least a subset of POA_{iso} neurons project to the caudolateral PAG where 409 neurons important for USV production reside, and an attractive possibility is that POA_{iso} 410 neurons promote USV production via disinhibition of PAG-USV neurons as previously 411 demonstrated for genetically-defined subsets of POA neurons (Chen et al., 2021; Michael et 412 al., 2020; Tschida et al., 2019). Future experiments will be required to determine whether 413 PAG-projecting POA_{iso} neurons also regulate social investigation and mounting, or 414 alternatively, whether distinct molecularly-defined or projection-defined subsets of POA_{iso} 415 neurons differentially regulate these different aspects of female social behavior. The latter 416 organization would be reminiscent of how projection-defined subsets of galanin-expressing 417 POA neurons regulate different aspects of parental behavior (Kohl et al., 2018).

Although reversible inhibition of POA_{iso} neurons reduced both social investigation and USV production in single-housed females, permanent caspase-mediated ablation of these neurons significantly reduced mounting but had no effects on social investigation or USV production. These differences are largely consistent with prior studies that reported effects on both mounting and USV production following reversible manipulations of POA activity and effects on mounting but not on USV production following irreversible manipulations of POA activity (Bean et al., 1981; Chen et al., 2021; Gao et al., 2019; Karigo et al., 2021; Wei et al., 425 2018). One possibility is that POA_{iso} neurons do not directly regulate USV production but 426 rather that reversible silencing of these neurons causes off-target disruptions of neural 427 activity in interconnected brain regions that in turn directly regulate USV production. Such a 428 relationship was demonstrated for motor cortex, whereby reversible silencing of motor cortex 429 disrupted performance of a learned forelimb reaching task in rats, while permanent lesions 430 of motor cortex had no effect on task performance after learning (Otchy et al., 2015). 431 However, our finding that optogenetic activation of POA_{iso} neurons elicits USV production, 432 along with past work demonstrating that POA activation elicits USV production (Chen et al., 433 2021; Gao et al., 2019; Karigo et al., 2021; Michael et al., 2020) supports the idea that the 434 POA directly regulates USV production. An alternative explanation for the apparently 435 contradictory effects of chemogenetic silencing vs. ablation of POA_{iso} neurons is that while 436 the POA plays an obligatory role in mounting behavior (at least in certain contexts, see 437 below), compensatory changes in additional, non-POA circuits that promote USV production 438 (and social investigation) are sufficient to compensate for the permanent loss of POA_{iso} 439 neurons, leaving these behaviors unperturbed following POA_{iso} neuronal ablation. The 440 identification of forebrain-to-midbrain circuits that regulate USV production in both females 441 and males remains an important future goal.

442 Our finding that POA_{iso} neurons regulate female-female mounting extends recent 443 work examining the role of hypothalamic regions in regulating male mounting behavior in 444 different social contexts (Karigo et al., 2021). Although the POA plays a well-established role 445 in the regulation of mounting behavior by male rodents during interactions with females, 446 interactions which are typically marked by high rates of USV production and are considered 447 affiliative, the authors found that the VMH regulates male mounting behavior during 448 interactions with other males, which is typically not accompanied by USV production and 449 often precedes fighting. Taken together with the findings of the current study, we conclude 450 that the POA regulates mounting behavior in both male and female mice that is directed 451 toward female social partners and is accompanied by USV production. The question of 452 whether the social behaviors exhibited by single-housed female mice during same-sex 453 interactions are indeed affiliative and how these behaviors shape future interactions with 454 female social partners are important questions that remain to be addressed. Although 455 chemogenetic silencing of POA_{iso} neurons tended to reduce female mounting and ablation of 456 POA_{iso} neurons significantly reduced female mounting, we were not able to reliably elicit 457 mounting behavior through optogenetic activation of POA_{iso} neurons (optogenetically-elicited 458 mounting was observed in n = 3 laser stimulations across N = 2 POA_{iso}-ChR2 females). We 459 note that previous studies that elicited mounting in male mice via optogenetic activation of 460 either Esr1⁺ POA neurons or POA neurons that co-express Esr1 and VGAT used longer

461 laser stimulation trains (15-30s) than we employed in the current study (Karigo et al., 2021; 462 Wei et al., 2018). It is possible that longer periods of optogenetic activation of POA_{iso} 463 neurons would be more effective in eliciting mounting in group-housed females, and another 464 possibility is that POA_{iso} neurons overlap somewhat but not perfectly with the genetically-465 defined groups of POA neurons manipulated in those prior studies. Regardless, our finding 466 that optogenetic activation of POA_{iso} neurons promotes USV production efficaciously relative 467 to mounting indicates that effects on USV production are not secondary to effects on other 468 female social behaviors.

469 To understand whether a similar population of POA neurons might regulate changes 470 in social behavior following short-term isolation in males, we extended our experiments to 471 single-housed males that subsequently interacted with either a male social partner or with a 472 female social partner. Consistent with our prior work (Zhao et al., 2021), we found that short-473 term isolation exerts more robust on the effects of male behavior during subsequent 474 interactions with females than during interactions with males. Although single-housed males 475 exhibited increased rates of social investigation while interacting with both males and 476 females relative to group-housed males, single-housed males that interacted with females 477 exhibited increases in all measured social behaviors (social investigation, USV production, 478 and mounting), a pattern of behavioral changes that is similar to what we observed in single-479 housed females. We also observed that a population of POA neurons increased Fos 480 expression in single-housed males following interactions with female visitors but not 481 following interactions with male visitors. Unexpectedly, we found that the effects of 482 chemogenetically inhibiting POA_{iso} neurons in males differed from those in females. Namely, 483 while reversible inhibition of POA_{iso} neurons tended to reduce mounting during interactions 484 between single-housed subject males and female visitors, there were no effects on rates of 485 social investigation and USV production. These findings differ also from recent work showing 486 that chemogenetic inhibition (Chen et al., 2021) or optogenetic inhibition (Karigo et al., 2021) 487 of Esr1⁺ POA neurons reduces USV production in male mice. Although the factors that 488 account for this difference in results are unclear, one possibility is that our TRAP2-based 489 viral labeling in males is biased toward POA neurons that regulate mounting as compared to 490 POA neurons that regulate USV production, although as stated above, more work is 491 required to understand whether these different social behaviors are regulated by distinct or 492 overlapping subsets of POA neurons. Future experiments can also explore to what extent 493 female and male POA_{iso} neurons represent molecularly, anatomically, and functionally 494 similar or dissimilar neuronal populations.

495 The current study adds to an emerging body of literature implicating the POA in the 496 regulation of social behavior in both females and males, including in non-sexual social 497 contexts (Fukumitsu et al., 2022; Hu et al., 2021; Liu et al., 2023; McHenry et al., 2017; Wu 498 et al., 2021). Our findings also complement recent work that has described a role for the 499 POA in regulating behavioral responses to social isolation in female mice. A recent study 500 using BALB/c mice found that reunion with same-sex cagemates following short-term 501 "somatic isolation" (i.e., separation from cagemates via a partition that permits visual, 502 auditory, and olfactory signaling but prevents physical contact) increases the activity of 503 calcitonin-receptor (Calcr) expressing POA neurons (Fukumitsu et al., 2023). Knockdown of 504 Calcr expression in these neurons reduced social-seeking behaviors directed at the cage 505 partition exhibited by single-housed females, and chemogenetic activation of these neurons 506 increased partition-biting behavior in single-housed females. Another recent study using 507 female FVB/NJ mice applied a TRAP2-based intersectional approach to identify POA neurons that increase their activity following reunion with same-sex cagemates (MPN_{reunion} 508 509 neurons) (Liu et al., 2023). Optogenetic activation of these neurons did not alter social 510 investigation in group-housed females, whereas activation of these neurons during reunion 511 with cagemates following short-term social isolation decreased social investigation. 512 However, optogenetic inhibition of MPN_{reunion} during reunion did not alter rates of social 513 investigation in single-housed females. Although we did not test the effects of activating 514 POA_{iso} neurons in single-housed females in the current study, the difference in the effects of 515 inhibiting POA_{iso} neurons on the behavior of single-housed females (reduced rates of social 516 investigation, USV production, and mounting) relative to the effects of optogenetic inhibition 517 of MPN_{reunion} neurons in Liu et al. (no effect on social investigation during reunion) suggest 518 that these may represent distinct populations of POA neurons. We note that in the current 519 study, single-housed female subjects remained single-housed for 24 hours following social 520 interaction TRAPing sessions, a design intended to maximize viral labeling of POA neurons 521 that promote increased female social behaviors and that would likely in turn minimize viral 522 labeling of POA neurons that promote social satiety. Taken together, these studies highlight 523 a complex role for the POA in regulating multiple aspects of changes in social behavior 524 following short-term social isolation. Although some studies of social isolation have avoided 525 the use of C57BL/6J female mice, the robust triad of changes in social behavior exhibited by 526 these females following short-term isolation affords a powerful opportunity to continue 527 investigating neural circuit mechanisms through which short-term social isolation promotes 528 social behaviors, as well as to investigate how hypothalamic circuits regulate the 529 coordinated production of suites of social behaviors during female-female social interactions.

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- 538
- 539 **Declaration of Interests**: The authors declare no competing interests.

540 Figure legends

541 Figure 1. The POA contains neurons that increase Fos expression in single-housed

542 females following same-sex interactions. (A) Schematic of experiment to measure Fos 543 expression in group-housed and single-housed females following same-sex social 544 interactions. (B) Total time spent engaged in resident-initiated social investigation for group-545 housed residents (teal) and single-housed residents (maroon). (C) Same as (B), for total 546 resident-initiated mounting time. (D) Same as (B), for total USVs recorded from pairs 547 containing group-housed or single-housed residents. (E) Left-most image shows the location 548 of the POA in a coronal brain section. Representative confocal images show Fos expression 549 (green) in the POA of a group-housed female (left) and a single-housed female (right) 550 following same-sex social interactions. Blue, Neurotrace. (F) Quantification of Fos-positive 551 neurons is shown for the POA (left), the VMH (middle), and the caudolateral PAG (right) for 552 group-housed and single-housed females. Open bars show data from females that did not 553 engage in social interactions with novel females (baseline), and closed bars show data from 554 females following social interactions with novel females (interaction). (G) Total time spent in 555 resident-initiated interaction is plotted for 14-day single-housed (maroon) and re-group-556 housed females (teal) during same-sex interactions that occurred prior to isolation (day 0), 557 following 3 days of isolation (day 3), and on the test day (day 17). (H) Same as (G), for total 558 resident-initiated mounting time. (I) Same as (G), for total USVs. (J) Quantification of Fos-559 positive POA neurons is shown for 14-day single-housed females (maroon) and re-group-560 housed females (teal).

561 Figure 2. Effects of chemogenetic inhibition of POA_{iso} neurons on the social behaviors

562 of single-housed female mice. (A) Experimental timeline and viral strategy to

- 563 chemogenetically inhibit the activity of POA_{iso} neurons in single-housed females. (B) Total
- time spent in resident-initiated social investigation is shown on saline and CNO days for 4
- 565 experimental groups: (red symbols) experimental females in which hM4Di is expressed in
- 566 POA_{iso} neurons; (black symbols) control females in which GFP is expressed in POA_{iso}
- 567 neurons; (brown symbols) control females in which hM4Di is expressed in 'TRAPed' AH
- neurons; (gray symbols) control females in which hM4Di is expressed in 'TRAPed' VMH
- neurons. (C) Same as (B), for total duration of resident-initiated mounting. (D) Same as (B),
- 570 for total USVs. (E) Total movement is plotted for females with hM4Di expressed in POA_{iso}
- 571 neurons, on saline days vs. CNO days.

572 Figure 3. Effects of caspase-mediated ablation of POA_{iso} neurons on the social

- 573 behaviors of single-housed female mice. (A) Experimental timeline and viral strategy for
- 574 caspase-mediated ablation of POA_{iso} neurons in single-housed females. (B) Total time spent

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575 in resident-initiated social investigation is shown pre- and post-4-OHT treatment for 2

576 experimental groups: (red symbols) experimental females in which caspase is expressed in

577 POA_{iso} neurons; (black symbols) control females in which GFP is expressed in POA_{iso}

neurons. (C) Same as (B), for total duration of resident-initiated mounting. (D) Same as (B),

579 for total USVs.

580 Figure 4. Effects of optogenetic activation of POA_{iso} neurons on the behaviors of

group-housed female mice. (A) Experimental timeline and viral strategy to optogenetically
 activate POA_{iso} neurons in group-housed females. (B) Mean total USVs produced during

583 social sessions shown for pre-laser periods and during laser stimulation periods for

584 experimental females with ChR2 expressed in POA_{iso} neurons (red symbols; N = 9) and for

585 control females with GFP expressed in POA_{iso} neurons (black symbols, N = 6). (C) Same as

586 (B), for social sessions. (D) Spectrograms are shown from a representative POA_{iso}-ChR2

587 female to illustrate USVs that were elicited through optogenetic activation of POA_{iso} neurons

588 in a solo session (top) and a social session (bottom). Blue bars indicate timing of laser

589 stimulation.

590 Figure 5. Context-dependent differences in POA Fos expression and effects of

591 chemogenetic inhibition of POA_{iso} neurons on the social behaviors of single-housed

592 male mice. (A) Total time spent in resident-initiated social investigation is shown for group-

593 housed male residents (teal) and single-housed male residents (maroon) during interactions

with either female visitors (left) or male visitors (right). (B) Same as (A), for total resident-

595 initiated mounting time. (C) Same as (A), for total USVs recorded from pairs containing

596 group-housed or single-housed male residents. (D) Total number of Fos-positive POA

597 neurons is shown for group-housed male residents (teal) and single-housed male residents

598 (maroon) following interactions with female visitors (left) or male visitors (right). (E)

599 Experimental timeline and viral strategy to chemogenetically inhibit the activity of POA_{iso}

600 neurons in single-housed males. (F) Total time spent in resident-initiated social investigation

601 is shown on saline and CNO days for experimental males in which hM4Di is expressed in

602 POA_{iso} neurons (red symbols, N = 7) and control males in which GFP is expressed in POA_{iso}

neurons (black symbols, N = 7). (G) Same as (F), for total duration of resident-initiated

604 mounting. (H) Same as (F), for total USVs.

605 Materials and methods

606 Key Resources

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
c-Fos (9F6) Rabbit mAb	Cell Signaling Technology	2250S
Alexa Fluor 488 goat-anti-rabbit	Invitrogen	A-11008
Neurotrace 435/455 Blue	Thermo Fischer Scientific	CAT#: N21479
Commercial assay or kit		
HCR v3.0	Molecular Instruments	
Bacterial and Virus Strains		
AAV2/1-hSyn-FLEX-hM4Di-mCherry	Addgene	RRID: Addgene_44262
AAV2/1-Ef1alpha-hChR2(h134R)-EYFP- WPRE-HGHpA	Addgene	RRID: Addgene_20298
AAVrg-pgk-Cre	Addgene	RRID: Addgene_24593
pAAV-flex-taCasp3-TEVp	Addgene	RRID: Addgene_45580
AAV2/1-pCAG-FLEX-EGFP-WPRE	Addgene	RRID: Addgene_51502
Chemicals, Peptides, and Recombinant P	roteins	
Clozapine N-oxide dihydrochloride	Hello Bio	HB6149; E1458-2-1
4-Hydroxytamoxifen	Hello Bio	HB6040
Experimental Models: Organisms/Strains		·
TRAP2 Fos ^{tm2.1(ICTP/ER12)LU0} /J	Jackson Labs	IMSR_JAX:030323
Ai14 B6.Cg <i>Gt(ROSA)</i> 26Sor ^{tm14(CAG-}	Jackson Labs	IMSR_JAX:007914
C57BL/6J	Jackson Labs	IMSR_JAX:000664
Software and Algorithms		·
MATLAB	Mathworks	http://www.mathworks.com RRID: SCR_001622
ZEN	Zeiss	https://www.zeiss.com RRID:SCR_013672
Spike2	CED	http://ced.co.uk RRID:SCR_000903
ImageJ	NIH	https://imagej.net/ij/ RRID:SCR_003070
Behavioral Observation Research Interactive Software	Open Behavior	https://github.com/olivierfriard/BORIS RRID:SCR_021509
R Project for Statistical Computing	R Core Team	http://www.r-project.org/ RRID:SCR_001905
R Studio	Posit	https://posit.co/ RRID:SCR_000432

607

608 Lead contact

609 Further information and requests for resources should be directed to and will be fulfilled by

610 the lead contact, Katherine Tschida (kat227@cornell.edu).

611 Experimental models and subject details

612 Animal statement

- 613 All experiments and procedures were conducted according to protocols approved by the
- 614 Cornell University Institutional Animal Care and Use Committee (protocol #2020-001).

615 Animals

616 TRAP2 (Jackson Laboratories, 030323) and Ai14 (Jackson Laboratories, 007914) mice were

- at least 8 weeks old at the time of the experiments or surgeries. TRAP2;Ai14 mice were
- 618 generated by crossing TRAP2 with Ai14. All mice were kept on a 12:12 reversed light/dark
- 619 cycle, were housed in ventilated micro-isolator cages in a controlled environment with
- 620 regulated temperature and humidity, and were provided with unrestricted access to food and
- 621 water. A running wheel (Innovive) was present in all homecages from the time of weaning
- and was subsequently removed immediately before initiating the social interaction test.
- Mouse cages were cleaned weekly, and experiments were never conducted on cage changedays.

625 Methods details

626 **Social isolation and social interaction tests**

627 Female and male subject mice were either group-housed with same-sex siblings or 628 separated from their cage mates and individually housed in clean cages for three days prior 629 to behavioral tests. In the case of group-housed subject mice, siblings were temporarily 630 removed from the home cage for the duration of the test. The subject animal's home cage 631 was then placed in a sound-attenuating recording chamber (Med Associates) equipped with 632 an ultrasonic microphone (Avisoft), an infrared light source (Tendelux), and a webcam 633 (Logitech, with the infrared filter removed to enable video recording under infrared lighting 634 conditions). A novel, group-housed visitor mouse (female or male mouse on a C57BL/6 635 background) was placed in the home cage of the subject mouse, and vocal and non-vocal 636 behaviors were recorded for 30 minutes. Visitor mice were used across multiple experiments 637 (< 6 in total), including in interactions with both group-housed and single-housed subject 638 mice. Visitor females used in male-female interactions were never used for female-female 639 experiments, but a subset of visitors used in female-female interactions were subsequently 640 used in male-female experiments.

A separate cohort of female mice was used to investigate the effects on social behavior of re-group-housing following a period of social isolation. For this cohort of mice, social

- 643 interaction tests with novel, group-housed female visitors were conducted at three
- timepoints: (1) on day 0, when subject females were still group-housed; (2) On day 3, after
- being single-housed for 3 days; (3) on day 17, after a randomly selected subset of subject
- 646 females were re-group-housed with their siblings for two weeks, and the remaining female
- 647 subjects remained single-housed for two weeks.

648 USV recording and detection

- 649 USVs were recorded with an ultrasonic microphone (Avisoft, CMPA/CM16), amplified
- 650 (Presonus TubePreV2), and digitized at 250 kHz (Avisoft UltrasoundGate 166H or CED
- Power 1401). USVs were detected with custom MATLAB codes (Tschida et al., 2019) using
- the following parameters (mean frequency > 45 kHz; spectral purity > 0.3; spectral
- discontinuity < 1.00; minimum USV duration = 5 ms; minimum inter-syllable interval = 30
- 654 ms).

655 Analyses of non-vocal social behaviors

- Trained observers used BORIS software (v.8.13; Friard and Gamba, 2016) to score the
- 657 following non-vocal behaviors: resident-initiated social investigation and resident-initiated
- mounting. Social investigation included sniffing and following. Resident-initiated mounting of
- the visitor typically occurred following a period of resident-initiated social investigation, with
- the resident mouse positioning its forelimbs on top of the body of the visitor, sometimes with
- 661 pelvic thrusts and sometimes without. Neither visitor-initiated mounting nor fighting were
- observed in our dataset.
- 663 In some trials, total movement was estimated using a custom MATLAB code that allows the
- user to mark the position of a mouse in every 30th frame (i.e., once per second). Total
- movement was then calculated as the sum of changes in position across pairs of markedframes.

667 Fos immunohistochemistry

- Two hours following the start of the social interaction test, mice were deeply anesthetized
- using isoflurane and then transcardially perfused with phosphate-buffered saline (PBS, pH
- 670 7.4), followed by 4% paraformaldehyde (PFA; Sigma-Aldrich, in 0.1 M PBS, pH 7.4). Brains
- 671 were subsequently dissected and post-fixed in 4% PFA for 24 hours at 4°C, followed by
- 672 immersion in 30% sucrose solution in PBS for 48 hours at 4°C. Afterward, brains were
- 673 embedded in frozen section embedding medium (Surgipath, VWR), flash frozen in a dry ice-
- ethanol (100%) bath, and then stored at –80°C until sectioning. Sections were cut on a
- 675 cryostat (Leica CM1950) to a thickness of 80 μm, washed in PBS (3 x 5 mins at RT),

676 permeabilized for 2-3 hours in PBS containing 1% Triton X-100 (PBST), and then blocked in

- 677 0.3% PBST containing 10% Blocking One (Nacalai USA) for 1 hour at RT on a shaker.
- 678 Sections were then incubated for 24 hours at 4°C with primary antibody in blocking solution
- 679 (1:1000 rabbit-anti-Fos, Cell Signaling Technologies, 2250S), washed 3 x 30 minutes in
- 680 0.3% PBST, then incubated for 24 hours at 4°C with secondary antibody in blocking solution
- 681 (1:1000, Alexa Fluor 488 goat-anti-rabbit, Invitrogen, plus 1:500 NeuroTrace, Invitrogen)
- Finally, sections were washed for 2 x 10 minutes in 0.3% PBST, followed by washing for 2 x
- 683 10 minutes in PBS. After mounting on slides, sections were dried and coverslipped with
- Fluromount G (Southern Biotech). Slides were imaged with a 10x objective on a Zeiss 900
- laser scanning confocal microscope, and Fos-positive neurons within regions of interest
- 686 were counted manually by trained observers.

687 Floating section two-color in situ hybridization

688 In situ hybridization was conducted using hybridization chain reaction (HCR v3.0, Molecular 689 Instruments). Ten minutes after the completion of the 30-minute social interaction tests, mice 690 underwent transcardial perfusion with RNase-free PBS (DEPC-treated), followed by 4% 691 PFA. Dissected brain samples were post-fixed overnight in 4% PFA at 4°C, cryoprotected in 692 a 30% sucrose solution in DEPC-PBS at 4°C for 48 hours, flash frozen in section embedding 693 medium, and stored at -80°C until sectioning. 40 µm-thick coronal floating sections were 694 collected into sterile 24-well plates in DEPC-PBS. These sections were briefly fixed once 695 again for 5 minutes in 4% PFA and subsequently immersed in 70% EtOH in DEPC-PBS 696 overnight. Sections were then rinsed in DEPC-PBS, incubated for 45 minutes in 5% SDS in 697 DEPC-PBS, followed by a series of rinses and incubations: 2x SSCT, pre-incubation in HCR 698 hybridization buffer at 37°C, and incubation in HCR hybridization buffer containing RNA 699 probes (VGAT and Fos) overnight at 37°C. Sections were then rinsed 4 x 15 minutes at 700 37°C in HCR probe wash buffer, rinsed in 2X SSCT, pre-incubated in HCR amplification 701 buffer, and then incubated in HCR amplification buffer containing HCR amplifiers at RT for 702 approximately 48 hours. On the final day, sections were rinsed in 2x SSCT, counterstained 703 with DAPI (Thermo Fisher, 1:5000), rinsed again with 2x SSCT, mounted on slides, and 704 coverslipped with Fluoromount-G (Southern Biotech). After drying, slides were imaged with a 705 10x or 20x objective on a Zeiss 900 laser scanning microscope. Neurons were scored from 706 three sections of tissue from the POA from each mouse, and the absence of presence of 707 staining for different probes was quantified manually by trained scorers.

708 Viruses

- 709 The following viruses and injection volumes were used: AAV2/1-hSyn-FLEX-hM4Di-mCherry
- 710 (Addgene #44262, 200 nL), AAV2/1-CAG-FLEX-EGFP-WPRE (Addgene #51502, 200 nL),

- 711 AAV2/5-Ef1alpha-FLEX-taCasp3-TEVp (Addgene #45580, 200 nL), AAV2/1-Ef1alpha-
- 712 hChR2(h134R)-EYFP-WPRE-HGHpA (Addgene #20298, 200 nL), AAVrg-pgk-Cre (Addgene
- 713 #24593, 200 nL). The final injection coordinates were as follows: POA, AP = -0.1 mm, ML =
- 0.6 mm, DV = 5.1 mm; AH, AP = -0.7 mm, ML = 0.6 mm, DV = 5.1 mm; VMH, AP = -1.5 mm,
- 715 ML = 0.7 mm, DV = 5.4 mm; PAG, AP = -4.7 mm, ML = 0.6 mm, DV = 1.75 mm. Viruses
- 716 were pressure-injected using a pulled glass pipettes mounted in a programmable nanoliter
- 717 injector (Nanoject III, Drummond) at a rate of 15 nL every 60 s.

718 Stereotaxic Surgery

- 719 Mice were anesthetized using isoflurane (2.5% for induction, then 1.5 2.5% for
- 720 maintenance) and then securely positioned in a stereotaxic apparatus (Angle Two, Leica). A
- 721 midline incision in the scalp was made to expose the skull, and small craniotomies were
- 722 created dorsal to each injection site using a surgical drill. Viral injection pipettes were left in
- 723 place for a minimum of 10 minutes before and after viral injections to minimize backflow
- vupon pipette withdrawal from the brain. Surgical sutures (LOOK 774B, Fisher Scientific) and
- tissue adhesive (3M) were used to close the incision.
- 726 For optogenetic activation experiments, an optogenetic ferrule (RWD Fiber Optic Cannula,
- 727 Ø1.25 mm Ceramic Ferrule, 200 μm Core, 0.22 NA, L = 7 mm) was implanted approximately
- 250 µm above the viral injection site immediately following the viral injection and was
- 729 secured in place with Metabond (Parkell).

730 TRAP activity-dependent labeling

- 731 Solutions of 4-hydroxytamoxifen (4-OHT, HelloBio, HB6040) were prepared by dissolving 4-
- OHT powder at 20 mg/mL in ethanol by shaking at 37°C, and aliquots (75 uL) were then
- stored at -20°C. Before use, 4-OHT was redissolved in ethanol by shaking at 37°C and
- filtered corn oil was added (150 uL). Ethanol was then evaporated by vacuum under
- centrifugation to give a final concentration of 10 mg/mL, and the 4-OHT solution was used
- on the same day it was prepared.
- 737 To express viral transgenes in recently active neurons, we used the Targeted
- 738 Recombination in Active Populations (TRAP2) strategy. Two weeks following viral injection,
- 739 TRAP2 and TRAP2;Ai14 mice were given 30-minute social encounters (as described
- above). Following the social encounter, subject mice received I.P. injections of 4-OHT (50
- 741 mg/kg) to enable expression of viral transgenes in recently active neurons. To minimize
- neural activity triggered by stimuli outside of the social interaction test, all subject animals

743 were individually housed for an additional 24 hours following 4-OHT treatment before being

744 re-group-housed with their same-sex siblings.

745 Chemogenetic inhibition

746 To reversibly reduce neuronal activity, TRAP2 female mice received bilateral injections of an 747 Cre-dependent inhibitory DREADDs virus into the hypothalamus (AAV2/1-hSyn-FLEX-748 hM4Di-mCherry; injected into the POA, AH, or VMH) as described above. TRAP2 male mice 749 received the same viral injections into the POA only. Three weeks later, mice were single-750 housed for 3 days and then were subsequently given a 30-minute social encounter with a 751 novel, group-housed female visitor. Subject mice then received an I.P. injection of 4-OHT to 752 enable expression of hM4Di in activity-defined populations of hypothalamic neurons. Two 753 weeks later, subject mice received an I.P. injection of either sterile saline (as a control) or 754 clozapine-n-oxide (CNO, 4 mg/kg, Hello Bio HB6149; to inhibit neurons expressing hM4Di) 755 30 minutes prior to a social interaction test. Three days later, mice that previously were 756 treated with saline received an I.P. injection of CNO, and mice that were previously treated 757 with CNO received an I.P. injection of saline, 30 minutes prior to another social interaction. 758 Rates of USV production and non-vocal social behaviors were compared between saline 759 and CNO days within animals to assess the effects of neuronal inhibition on social 760 behaviors. Control mice received unilateral injections into the POA of a Cre-dependent AAV 761 driving the expression of GFP (AAV2/1-CAG-FLEX-EGFP) and were otherwise treated

762 identically to experimental animals.

763 Neuronal ablation

To permanently ablate neurons, TRAP2;Ai14 female mice received bilateral injections of an
 AAV2/5-ef1alpha-FLEX-taCasp3-TEVp virus into the POA. Following a three-week recovery

- period, these animals were individually housed for three days and subsequently given a 30-
- 767 minute social encounter with a novel, group-housed female visitor. Subject mice then
- received an I.P. injection of 4-OHT to enable expression of caspase in activity-defined POA
- neurons. Two weeks later, females were single-housed for three days and then given a
- second 30-minute social interaction test. Social behaviors of subject females were compared
- between the pre-ablation and post-ablation interaction tests to assess the effects of neuronalablation.

773 Optogenetic activation

- 774 Female TRAP2 mice received unilateral injections into the POA of AAV-ef1α-FLEX-ChR2
- 775 (experimental) or AAV-CAG-FLEX-GFP (control). In the same surgery, an optogenetic

776 ferrule was implanted approximately 250 µm above the viral injection site. Three weeks later, 777 females were single-housed for 3 days and then given a 30-minute social interaction with a 778 novel group-housed, female visitor. Subject females then received an I.P. injection of 4-779 OHT. Two weeks later, females were first placed alone in the testing chamber for a 5-minute 780 habituation period after connecting the laser patch cable to the female's optogenetic ferrule. 781 Optogenetic activation sessions consisted of a 5-minute period in which optogenetic 782 activation was performed in solo females, followed by a 20-minute period in which activation 783 was performed as subject females interacted with a novel, group-housed female visitor. The 784 social session was further divided into three phases: 5 minutes without optogenetic 785 activation, 10 minutes with optogenetic activation, and 5 minutes without optogenetic 786 activation. During the middle 10 minutes of the social session, some laser stimuli were 787 delivered at times when the two females were near to one another (inter-animal distance < 788 ~2 mouse body lengths), and other stimuli were delivered at times when the females were 789 not in close contact. POA_{iso} neurons were optogenetically activated with illumination from a 790 473 nM laser (10 mW) at 20-50 Hz (10-20 ms pulses, trains lasted 5-10s) Laser stimuli were 791 driven by computer-controlled voltage pulses (Spike 2 version 10.8, CED).

792 Anatomical tracing

793 Female TRAP2 mice used as GFP controls in the chemogenetic inhibition experiments were 794 subsequently used for anterograde mapping of the axonal projections of POA_{iso} neurons. 795 Three weeks following unilateral injection into the POA of a Cre-dependent AAV driving the 796 expression of GFP (AAV2/1-CAG-FLEX-EGFP), females were single-housed for 3 days and 797 subsequently given a 30-minute social interaction test. Subject mice then received an I.P. 798 injection of 4-OHT. Six weeks later, females were perfused, brains were collected and 799 sectioned, and a confocal microscope was used to image GFP-positive axon terminals 800 within coronal brain tissue sections.

801 To examine the overlap between PAG-projecting POA neurons and Fos expression, Ai14 802 females first received a unilateral injection into the PAG of an AAV driving the retrograde 803 expression of Cre-recombinase (AAVrg-pgk-Cre). Two weeks later, these females were 804 given a 30-minute social interaction test. Ninety minutes after the test, subject females were 805 perfused, brains were collected, and coronal brain sections containing the POA were 806 collected for Fos immunohistochemistry as described above. Brain tissue sections were 807 imaged with a 10x objective on a Zeiss 900 laser scanning confocal microscope, and POA 808 neurons that were Fos-positive and tdTomato-positive were counted manually by trained 809 observers.

810 Statistics

- 811 Two-sided statistical comparisons were used in all analyses (alpha = 0.05). The Shapiro-
- 812 Wilk test was performed to analyze the normality of each data distribution, and non-
- 813 parametric statistical tests were used for comparisons that included non-normally distributed
- 814 data. No statistical methods were used to pre-determine sample size. Mice were only
- 815 excluded from analysis in cases in which viral injections were not targeted accurately.
- 816 Details of the statistical analyses used in this study are included in Table S1.

817

818 Supplemental Information

819 Figure S1. Additional characterization of Fos expression in single-housed vs. group-

820 housed females and comparison to rates of female social behaviors. (A) 821 Representative confocal images show Fos expression (green) in the VMH of a group-822 housed female (left) and a single-housed female (right) following same-sex social 823 interactions. Blue, Neurotrace. (B) Same as (A), for the caudolateral PAG. (C) Left, the 824 relationship between total time spent in resident-initiated social investigation and numbers of 825 Fos-positive POA neurons is shown for group-housed (teal) and single-housed (maroon) 826 female residents following interactions with novel females. Middle, same as left, for total 827 resident-initiated mounting time vs. numbers of Fos-positive POA neurons. Data only shown 828 for single-housed residents because group-housed residents never mounted female visitors. 829 Right, same as left, for total USVs vs. numbers of Fos-positive POA neurons. (D) Left, the 830 relationship between total time spent in resident-initiated social investigation and numbers of 831 Fos-positive VMH neurons is shown for group-housed (teal) and single-housed (maroon) 832 female residents following interactions with novel females. Middle, same as left, for total 833 resident-initiated mounting time vs. numbers of Fos-positive VMH neurons. Right, same as 834 left, for total USVs vs. numbers of Fos-positive VMH neurons. (E) Left, the relationship 835 between total time spent in resident-initiated social investigation and numbers of Fos-836 positive caudolateral PAG neurons is shown for group-housed (teal) and single-housed 837 (maroon) female residents following interactions with novel females. Middle, same as left, for 838 total resident-initiated mounting time vs. numbers of Fos-positive PAG neurons. Right, same 839 as left, for total USVs vs. numbers of Fos-positive PAG neurons.

840 Figure S2. Characterization of neurotransmitter phenotype and axonal projections of

841 **POA**_{iso} **neurons.** (A) Representative confocal images of in situ hybridization performed on

brain sections containing the POA, showing overlap of expression of Fos (green) and VGAT

843 (magenta). Blue, DAPI. (B) Quantification of proportion of Fos-positive POA neurons that

844 expressed VGAT. (C) Experimental timeline and viral strategy to express GFP in POA_{iso}

845 neurons. (D) Confocal images showing GFP-labeled axons of POA_{iso} neurons within the

846 caudolateral PAG. Blue, Neurotrace. (E) Experimental timeline and viral strategy to

- 847 retrogradely label PAG-projecting POA neurons with tdTomato. (F) Confocal image showing
- tdTomato labeling in a coronal section containing the POA, and dotted circles in insets
- 849 indicate examples of double-labeled neurons. Neurotrace, blue. (G) Quantification of
- proportion of tdTomato-expressing POA neurons that are also Fos-positive.

851 Table S1. Details of statistical analyses.

852

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Figure 1.

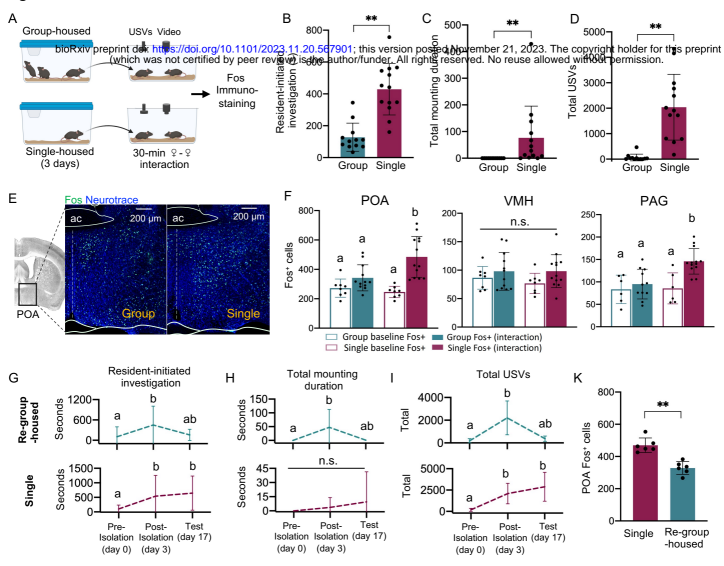


Figure 2.

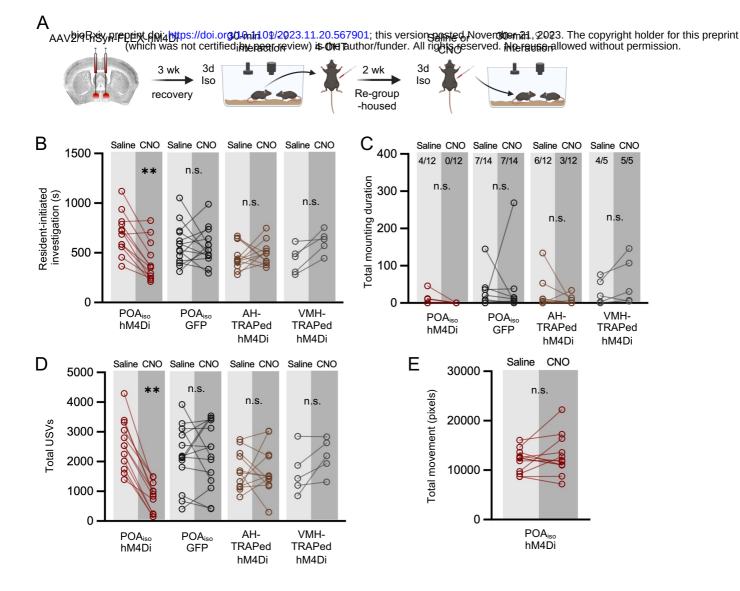


Figure 3.

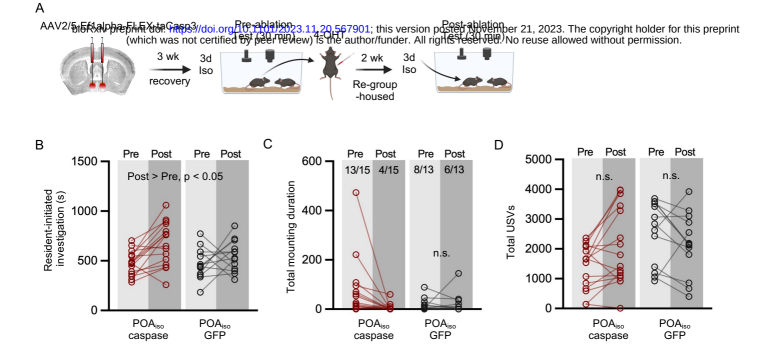


Figure 4

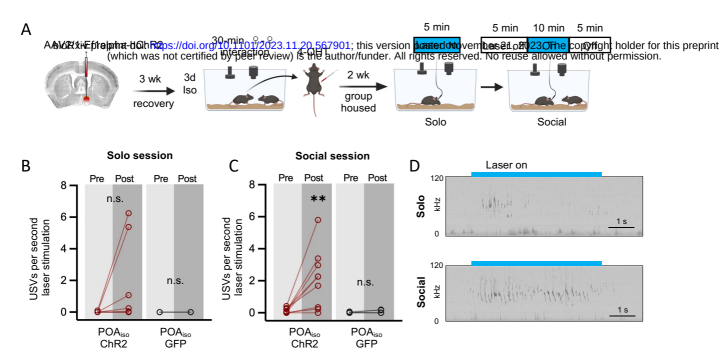


Figure 5.

