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Anxiolytic effects of Fluoxetine and Nicotine exposure on exploratory behavior in Zebrafish

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ABSTRACT

Zebrafish (Danio rerio) have emerged as a popular model for studying pharmacological effects on behavior 10 and anxiety. While there have been numerous studies documenting the anxiolytic and anxiogenic effects 11 of common drugs in zebrafish, many do not report or test for behavioral differences between the sexes. 12 Previous studies of zebrafish have indicated that males and females differ in their behavioral responses 13 to anxiety. In this study, we test for sex-dependent effects of fluoxetine and nicotine. We exposed fish to 14 system water (control), 10 mg/L fluoxetine, or 1 mg/L nicotine for three minutes prior to being subjected 15 to four minutes in an open-field drop test. Video recordings were tracked using ProAnalyst. Fish from 16 both drug treatments reduced swimming speed, increased vertical position, and increased use of the 17 top half of the open field when compared with the control, though fluoxetine had a larger effect on depth 18 related behaviors while nicotine mostly affected swimming speed. A significant sex effect was observed 19 where females swam at a slower and more constant speed than males in all treatments. No interactions 20 between sex and the drugs were observed across the entire study. 21

22 Keywords: behavior, anxiety, fluoxetine, nicotine

23 INTRODUCTION

The zebrafish (Danio rerio) is a popular research model for studying pharmacology (summarized in 24 Barros et al., 2008; Langheinrich, 2003) and behavior (Gerlai, 2015), particularly with regard to stress 25 and anxiety. The zebrafish provides a vertebrate model that breeds rapidly, is easy to maintain in large 26 numbers, and can be administered drugs through immersion. Zebrafish also share many of the same 27 neurotransmitters (Shin and Fishman, 2002) and stress pathways as humans, utilizing cortisol rather than 28 corticosteroids as used by rats and mice (Barcellos et al., 2007). These features have facilitated zebrafish 29 studies on addiction (Mathur and Guo, 2010), learning (Sison and Gerlai, 2010), social behavior (Buske 30 and Gerlai, 2014; Gerlai, 2014) and anxiety behaviors (Mathur and Guo, 2010; Maximino et al., 2010). 31 Anxiety-related behaviors are known to vary by sex in zebrafish and other model organisms, and these 32 differences may be explained by gonadal hormones(Zimmerberg and Farley, 1993; Palanza, 2001). Male 33 and female rats differ in their time spent in the center of an open field and a plus maze, though the nature 34 of these differences are also dependent on the strain observed (Mehta et al., 2013). In zebrafish, females 35 36 tend to be less anxious, or more bold, than males when measuring location preferences in the presence of a human observer (Benner et al., 2010; Oswald et al., 2013a). 37

Drugs are used to manipulate anxiety and related disorders in humans and are also a utilized as tool for understanding behavior. Fluoxetine, for example, is a drug used to treat depression and anxiety. It works by blocking the reuptake of serotonin in the brain (Beasley et al., 1992). Serotonin and its transporters have been associated with anxiety (Graeff et al., 1997; Lesch et al., 1996). Nicotine is naturally found in tobacco products and binds to nicotinic cholinergic receptors (nAChRs) to release dopamine (Benowitz

et al., 2009). The result is an anxiolytic response (Picciotto et al., 2002).

⁴⁴ Observations of male and female differences in anxiety-related behavior have led us to ask whether ⁴⁵ the effects of anxiolytic substances also differ by sex. There is evidence that the effectiveness of anxiolytic

drugs may vary with sex in mammals. Differential responses have been observed in humans utilizing 46 Sertraline, a selective serotonin reuptake inhibitor (SSRI) where females showed an enhanced response 47 compared to males. (Kornstein et al., 2000). Sex-specific differences were observed in the effectiveness 48 of the SSRI Fluoxetine in humans (Martényi et al., 2001), and studies utilizing rats (Mitic et al., 2013; 49 50 Leuner et al., 2004; Lifschytz et al., 2006) and mice (Monleón et al., 2002; Hodes et al., 2010) have shown a discrepancy between the sexes in both the physiological and behavioral responses to this drug where 51 efficacy tends to be greater in females than in males. Evidence in rats also suggest that nicotine's effects 52 on stress and anxiety may also differ between the sexes with males exhibiting a greater anxiolytic effect 53 (Faraday et al., 1999). This is important from a pharmacological standpoint in that effective doses may 54 differ between males and females. On a broader level, studies utilizing a single sex, or ignoring the effect 55

of sex altogether ought not to be used to draw broad conclusions about the effects of that drug.
 While zebrafish are becoming a model for pharmacological research, literature describing sex dependent effects of anxiolytic drugs in this system are lacking. In this experiment, we test the hypothesis

that zebrafish exhibit sex-dependent responses to fluoxetine and nicotine. These substances were chosen
 because they have known anxiolytic effects across a wide variety of model systems including humans

⁶¹ (Gilbert, 1979; Griffin and Mellon, 1999), rats (Cohen et al., 2009; Zhang et al., 2000) and zebrafish (Ben-⁶² can and Levin, 2008; Bencan et al., 2009; Cachat et al., 2010; Levin et al., 2007), and while sex-specific

effects have been observed in mammals, studies in zebrafish utilizing these substances largely ignore the

64 effects of sex.

METHODS

66 Subjects

67 Experimental fish were bred from adult Scientific Hatcheries strain (Huntingdon, CA) that has been maintained in our facility. Water in our Aquaneering Inc. (San Diego, CA) system was constantly 68 circulating and kept at a temperature of 28.5 °C on a 14 hour light:10 hour dark cycle. The fish were fed a 69 diet of brine shrimp twice and flake food (Tetramin) once for a total of three daily feedings. At the time of 70 data collection, the fish were four months old and housed in three-liter tanks in groups of five to achieve 71 maximal growth rates. Though zebrafish stocked at this density are known to develop social hierarchies 72 that can influence stress and behavior (Pavlidis et al., 2013), we randomly assigned individuals to a drug 73 treatment group such that these effects should be equally distributed across treatments. All aspects of 74 this study were approved by the University of Idaho's Animal Care and Use Committee under protocol 75 2014-14. 76

77 Dosing

Fluoxetine (generic (Teva Pharmaceuticals) from Wal Mart) and nicotine (Sigma Aldrich) treatments were 78 administered at concentrations of 10 mg/L for the fluoxetine and 1 mg/L for the nicotine. These doses 79 vary from standard doses in the zebrafish literature. Fluoxetine is often given at concentrations up to 100 80 μ g/L, but administered chronically over a two-week period (Egan et al., 2009). We used a higher dose 81 than the chronic concentrations reported in the literature, however it is important to note that this choice 82 could yield non-target effects due to higher concentrations. Nicotine is often administered as a ditartrate 83 salt at concentrations up to 100 mg/L (Levin et al., 2007). We used pure nicotine and were unsure at the 84 time of the experiment how the two forms compared with each other. We chose our dose based on the 85 LD50 concentration (4 mg/L) to avoid lethal effects on our subjects. Each drug was dissolved in system 86 water to make a working solution each morning of administration. A third treatment of only system 87 water served as a control. Fish were netted from their home tank and immediately placed into a beaker 88 containing 100mL of one of the three treatments. After three minutes of exposure to the drug dose, the 89 fish were transferred to an open field test tank filled with untreated system water for behavioral recording. 90 Dosing and behavioral observations were made on one fish at a time and the treatment type and order 91 were randomized across individuals. 92

Behavior Assay & Video tracking

⁹⁴ The fish were placed in a rectangular tank with interior dimensions measuring 25cm wide, 12 cm high

- 95 (from water level to bottom), and 6 cm thick (front to back). The volume of water in the tank was
- ⁹⁶ approximately 2 L. Each fish was filmed for four minutes (240 seconds) at 25 frames/second beginning
- ⁹⁷ from the time that the subject entered the water. The camera and operator were hidden behind a blind

during the recorded observation time. The tank was backlit with an opaque diffuser for the purposes of 98 creating a silhouetted object for motion tracking. After the four-minute period, the fish was netted out of 99 the test tank, placed into its own individual 1.5 L housing and returned to the main system. Observations 100 were recorded over three days between the hours of 10:00 am and 2:00 pm. After all subjects had been 101 recorded, weight and standard length measurements were obtained by first anesthetizing the individual in 102 MS-222 solution and blotting excess water with a paper towel. At this time, we also recorded the sex of 103 the individual using visual cues: larger, rounded abdomen and dull fins for females, smaller and leaner 104 abdomen and bright yellow fins for males. 105

Videos were digitally tracked using ProAnalyst[®] (Xcitex, Cambridge, MA). Tracking began with the 106 frame in which the fish hit the surface of the water, and proceeded to the end of the video. The tracking 107 data were imported into R for cleaning and processing. Each track was truncated to exclude the first 108 five seconds during which the fish would sink, but remain otherwise motionless, as it recovered from the 109 initial shock of being released from the net. Tracks were then standardized to 4 minutes, or 6000 frames. 110 We computed velocity from the x-y data points. Since the tracking software did not always track the exact 111 same position on the fish, velocity was estimated using the change in coordinates between two frames 112 before and two frames after the focal frame. This algorithm sufficiently smoothed the speed data while 113 retaining detail at small time intervals. 114

115 Analysis

116 Freezing

Freezing time was defined as the time a subject spent motionless on the bottom of the tank. We defined 117 motionless as maintaining a velocity of less than .01 cm/frame for more than 20 consecutive frames. Any 118 short bursts of motion flanked by considerable freezing times were verified in the video to be true motion. 119 If a time period of activity was less than 40 frames, it was re-categorized as part of the freezing time 120 as this motion is likely an artifact of the automated tracking. The freezing time was then calculated by 121 counting the total number of frames marked as frozen. We also characterized freezing behavior as a binary 122 'yes' or 'no' response as the propensity to show any freezing behavior can be considered an independent 123 response from duration of freezing. 124

125 Speed

We computed the average speed for each individual using only the active (non-frozen) data points from 126 the swim tracks. Freezing behaviors can cause a high degree of correlation with average swimming 127 behaviors such as speed and depth use. Since we analyzed freezing behavior separately, we chose to 128 analyze the effects of anxiolytic drugs on velocity during active swimming only. We predicted that anxious 129 individuals would swim slower on average than less anxious individuals (Gerlai et al., 2009). In addition, 130 we computed the variance in velocity for the active data points. The variance represents the consistency in 131 swim speed within an individual. Less anxious individuals should display more consistency in velocity 132 than more anxious individuals due to erratic behavior (Gerlai et al., 2009). 133

134 **Depth**

Depth was measured by the y-coordinate position in the swim track. We aligned the y origin with the 135 water's surface, and measured depth as increasing negatively toward the bottom of the tank. As with 136 velocity, depth variables were calculated using only the active points in the tracks. We analyzed both the 137 mean and variance (consistency) of depth. We predicted that anxious individuals should spend more time 138 near the bottom of the tank and should have a lower variance in depth (Levin et al., 2007; Oswald et al., 139 2013b). Conversely, we predicted that less anxious individuals will position themselves higher in the 140 water on average and spend more time exploring the entire tank, resulting in a larger variance in depth 141 usage. We also quantified at the number of times an individual entered the top half of the tank from the 142 bottom half. Such behavior may be indicative of anxiety, as anxious individuals tend to enter the top half 143 less often than less anxious individuals (Egan et al., 2009). We also expected that anxious individuals 144 would spend a smaller proportion of active swimming time in the top half, and that they would exhibit a 145 longer latency to enter the top from the beginning of the trial (Egan et al., 2009). The threshold between 146 the top and bottom halves was defined at -6 cm. 147

148 Horizontal Place Preference

¹⁴⁹ The width of the tank was divided into three equal sections and the proportion of time in the middle section

calculated to differentiate preference to be located in the center versus the edge of the test environment.

While we had clear expectations for location preference with respect to depth, it was unclear at the time of analysis whether the middle or the edges represent a "safe" zone with respect to horizontal preference. Experiments with rodents have found that stressed individuals prefer the edges of their arenas (thigmotaxis), but that this behavior is analogous to stressed fish preferring the bottom (Levin et al., 2007).

155 Statistical Analysis

We began with a MANOVA on all continuous variables where all individuals could be included. We 156 applied transformations where they were required to conform to the assumptions of normality in the 157 residuals (see Results for transformations). The initial model included the effects of weight as a covariate, 158 sex, drug treatment, and the sex by drug interaction. No significant effect of weight was observed, 159 and there was no improvement to the model by keeping the term, so we excluded weight from all 160 subsequent analyses. We performed individual ANOVAs on each of the continuous variables. Since 161 freezing occurrence is a binary response, it was analyzed using a logistic GLM to estimate and compare 162 the probability that an individual will freeze based on a given treatment group. In order to accurately assess 163 freezing time, only individuals that froze were used (N=52). All tests were performed with a significance 164 threshold of $\alpha = 0.05$. When a significant effect of drug treatment was detected, we performed pairwise 165 T-tests among the three treatments with a Tukey correction. 166

167 **RESULTS**

We recorded observations from 90 individuals divided equally and randomly among the 3 treatments (n=30 per treatment). Due to complications with the filming, observations on three of the individuals had to be removed leaving us with final sample size of 87 individuals broken down by treatment and sex as follows: 29 in the control treatment (17 females and 12 males), 30 in the fluoxetine treatment (16 females

and 14 males), and 28 in the nicotine treatment (14 females and 14 males).

173 Multivariate

The full model Type-II MANOVA included the effects of weight, sex, drug treatment, and the sex 174 by drug interaction on average depth, variance of depth, average speed, variance of speed, percent 175 of time spent in the top half, number of crosses into the top half, latency to enter the top half, and 176 proportion of time spent in the middle third horizontally (ie, away from the edges). There was a non-177 significant effect of sex ($\Lambda = 0.17896$, $F_{8.73} = 1.9889$, p = 0.05974) and a significant effect of drug 178 treatment ($\Lambda = 0.56646, F_{16,148} = 3.6551, p = 0.00001305$) on behavior, but no significant interaction. 179 There was no significant effect of weight as a co-variate, and including weight in the model showed no 180 improvement over removing it ($\Lambda = 0.95793, F_{5.76} = 0.66755, p = 0.6492$). With the reduced model, 181 we observed a significant effect of sex ($\Lambda = 0.22404, F_{8,74=2.6707}, p = 0.01237$) and drug treatment 182 $(\Lambda = 0.56659, F_{16,150} = 3.7057, p = 0.00001014)$. Therefore, for all subsequent analyses we considered 183 only the effects of sex, drug treatment, and the interaction term. 184

185 Individual components of behavior

We observed no significant interactions between sex and drug treatment in any of the individual behavior 186 components (see Table 1), consistent with the results of the MANOVA above. All components indicated a 187 significant effect of drug treatment (p < 0.05) except for freezing occurrence and freezing duration. The 188 subsequent descriptions describe the results of the post-hoc pairwise comparisons of the drug treatments 189 using the least-squared means and Tukey adjusted p-values based on 3 tests. We also observed a significant 190 effect of sex with regard to average swimming speed ($F_{1,81} = 10.7178, p = 0.001562$) and consistency 191 (variance) of swimming speed ($F_{1,81} = 13.9196, p = 0.0003528$). Males were on average faster than 192 females, but also exhibited less consistency in their swimming speeds. These were the only instances in 193 which the sexes differed in their behavior. 194

195 Freezing behavior

¹⁹⁶ Freezing behavior is a commonly observed anxiety related behavior in zebrafish (Egan et al., 2009). Of the

- ¹⁹⁷ 87 individuals observed, 52 exhibited freezing behavior. Though males tend to be more likely to freeze than
- females on average, this difference was not statistically significant ($\chi^2 = 3.7866, p = 0.05167$). We also
- failed to observe a significant effect of drug treatment on freezing occurrence ($\chi^2 = 3.7964, p = 0.14983$)
- as well as a sex by drug interaction ($\chi^2 = 0.3949, p = 0.82083$). For freezing duration, or latency to
- explore, we only included the 52 individuals that exhibited freezing behavior (control: F=11, M=10;

	Sex	Drug	Interaction	Fluoxetine	Nicotine
Freezing Time	0.17	0.26	0.76	0.99	0.33
Average Speed	0.00	0.01	0.63	0.13	0.00
Variance Speed	0.00	0.02	0.47	0.57	0.02
Average Depth	0.98	0.00	0.94	0.00	0.04
Variance Depth	0.62	0.01	0.91	0.01	0.19
Proportion in Top	0.86	0.00	0.72	0.00	0.22
Crosses to Top	0.57	0.00	0.89	0.00	0.45
Latency to Top	0.64	0.00	0.89	0.03	0.00
Proportion in Center	0.19	0.00	0.36	0.00	0.99

Table 1. Table of P-values summarizing results. **Bold** items are considered to show significant differences among treatment groups ($\alpha = 0.05$). P-values for the Fluoxetine and Nicotine columns represent pairwise comparisons with the control and are adjusted using the Tukey method for 3 comparisons.

fluoxetine: F=7, M=8; nicotine: F=6, M=10). This improved the assumptions of normality required for the ANOVA. Results of the type II ANOVA suggest that neither sex nor drug treatment have any significant effect on freezing duration (Sex: $F_{1,46} = 1.9604$, p = 0.1682; Drug: $F_{2,46} = 1.3707$, p = 0.2641). Figure 1 shows the results of freezing behaviors.

206 Speed

When analyzing only the active swimming data from the trials, fish from both drug treatments appear to reduce their average swimming speed compared with the control, however this pattern is only significant in the nicotine treatment (t = 3.373, p = 0.0032, see figure 2). Drugged fish also swam at a more consistent speed than the undrugged control fish ($F_{2,81} = 4.0654$, p = 0.0207731), but again this trend was only significant in the nicotine treatment (t = 2.818, p = 0.0166).

212 **Depth**

Both the subjects dosed with nicotine and fluoxetine positioned themselves higher in the water column than the control fish (nicotine: t = -2.462, p = 0.0417; fluoxetine: t = -4.711, p < .0001). Fish dosed with fluoxetine explored more of the water column than control subjects (t = -3.172, p = 0.0060). Subjects dosed with nicotine also exhibited more variation in depth use on average than the control subjects, but this difference was not significant (see figure 3).

We also divided the tank into two discrete and equal vertical zones and compared the proportion of time spent in the upper half (figure 4). Subjects dosed with fluoxetine tended to spend more than twice as much time in the upper half as control subjects and this difference is significant (t = -3.883, p = 0.0006). Subjects in both the nicotine and fluoxetine treatments exhibited a reduced latency time to first enter the top half than control subjects (nicotine: t = 3.333, p = 0.0037; fluoxetine: t = 2.652, p = 0.0258). When comparing the total number of visits to the top half, only the fluoxetine group showed a significant increase over the control (t = -3.801, p = 0.0008).

225 Horizontal Place Preference

All subjects spent most of their time near the edges avoiding the center (figure 4), consistent with the concept of thigmotaxis. However, subjects dosed with fluoxetine spent less time in the center and more time near the edges than subjects in the control and nicotine treatments (t = 3.257, p = 0.0046) which is inconsistent with a reduction in thigmotaxis resulting from a reduction in stress. At this time we are unsure how these results relate to anxiolytic properties of the drug.

231 DISCUSSION

232 Differences in fluoxetine and nicotine behavioral responses

Small prey fish such as zebrafish tend to behave in such a way as to reduce risk of predation. When placed

in a novel open field, such behavioral strategies include diving to the bottom and remaining motionless

- (Egan et al., 2009), and avoiding potentially risky locations such as the surface of the water (Wilson and
- Godin, 2009; Oswald et al., 2013b). Exposure to anxiolytic drugs alters these behaviors in ways that

may indicate an association between anxiety related behaviors and risk management. We observed a 237 decrease in bottom dwelling and an increase in time spent in the top half of the tank in fish exposed to 238 fluoxetine (figures 3 and 4). This is consistent with patterns observed by Egan et al. (2009) who also 239 report an increased use of the top of the water column by zebrafish exposed to fluoxetine. However, the 240 study by Egan et al. (2009) also reports a reduction in freezing bouts and freezing time, a pattern we 241 failed to observe. One explanation for this discrepancy could be differing effects of chronic and acute 242 dosing. Fluoxetine is metabolized into norfluoxetine, its active metabolite, in the liver by cytochrome 243 P450 enzymes (Rasmussen et al., 1995). It then travels through the bloodstream to the brain where it 244 blocks the reuptake of serotonin (Beasley et al., 1992). Metabolism of the drug could delay its effect 245 246 until after the animal had already recovered from freezing behavior. While most fluoxetine studies utilize chronic exposure, we have shown that similar behavioral changes can occur with just a single acute 247 dose. Acute exposure to fluoxetine has also been shown to reduce cortisol levels of zebrafish exposed 248 to a stressful environment (de Abreu et al., 2014). We speculate that the behaviors we observed may be 249 due to a reduction in physiological stress response resulting from exposure to the drugs, though more 250 experiments are needed to confirm this. 251

We observed changes in swimming speed, average depth, and latency to enter the top in fish exposed 252 to nicotine. Fish exposed to nicotine were quicker to enter the top and swam higher in the water column on 253 average compared to control fish. This is consistent with a reduction in anxiety related behaviors as seen 254 in the fluoxetine treatment group. Exposure to nicotine and fluoxetine appeared to decrease swimming 255 speed while increasing the consistency at which the fish swam. The increased consistency (reduction 256 of individual variance) might be explained by a reduction in anxiety, where individuals that are calm 257 should move at a fairly normal and constant pace, while anxious individuals may constantly alter their 258 swimming speeds in an erratic fashion Gerlai et al. (2009). Egan et al. (2009) reported an increase in 259 average swim speed with exposure to fluoxetine, which contrasts with our observations of slower average 260 swim speeds with exposure to either fluoxetine or nicotine. Sackerman et al. (2010) suggests that nicotine 261 may have sedating effects which could account for the slower swim speeds. However, we also observe 262 slower average swim speeds in the fluoxetine treatment, and though the difference is not statistically 263 different from the control, it is also not different from the nicotine effect. We observed a similar pattern in 264 the nicotine treatment with respect to the time spent at the top and the variation in depth use, where the 265 nicotine treatment was statistically indistinguishable from both the control and the fluoxetine treatments. 266 In these two instances, it is likely that the nicotine is having an anxiolytic effect, but that we used too low 267 of a dose to observe an effect that is different from the control. Sackerman et al. (2010) also failed to 268 observe an effect of nicotine on swim depth using a low dose of 25 mg/L, but noted that higher doses 269 such as 50 mg/L and 100 mg/L do produce a significant effect (Levin et al., 2007). Our dose of 1 mg/L 270 is noticeably lower than other studies of nicotine in adult Zebrafish, accounting for the our use of pure 271 nicotine liquid while the other studies used a nicotine tartrate salt (Levin et al., 2007; Sackerman et al., 272 2010). It should be noted that the relationship between the tartrate salt and pure form is about 0.325, such 273 that a concentration of 100mg/l of the tartrate equates to a concentration of 32.5mg/l of pure nicotine 274 (Matta et al., 2007). 275

Both nicotine and fluoxetine affected behavior in ways indicative of a reduction of anxiety. However, the two drugs also appear to affect different components of behavior. Nicotine had its highest effect on swimming speed, while fluoxetine mostly affected behaviors related to vertical positioning. This suggests that anxiety is not a simple condition, but rather a complex idea encompassing a number of components that are sometimes correlated, but not always connected. These behavioral components may be separated by different physiological pathways which could explain why different classes of drugs affect specific behaviors.

The effect of sex on behavior and drug efficacy

Sex differences in anxiety behaviors have been described in a number of species including rats (Mehta et al., 2013), stickleback (King et al., 2013), and guppies (Harris et al., 2010). While most of these studies find that males are typically more bold (less anxious) than females, our lab has previously observed the opposite trend in the Scientific Hatcheries strain of zebrafish with regard to association with humans, vertical position, and feeding latency in individual home-tank observations (Oswald et al., 2013a,b; Benner et al., 2010). These differences are the basis for our inquiry as to whether substances known to alter these behaviors might work at different efficacy in males and females. In the present study, we only observe

significant behavioral differences between the sexes with respect to swimming speed. While males swim slightly faster than females, it's the females that swim at a more constant rate. In addition, males seem to show a higher probability to exhibit freezing behavior across all three treatments, and even though this trend isn't statistically significant, it still leads us to suggest that males **could be** behaving with higher anxiety levels than females.

With the active swimming behaviors, we fail to observe differences between the sexes, and across all 296 of the behaviors, the data do not suggest any indications of sex-specific effects of either drug. There is 297 plenty of literature in mammalian models that contradict these findings (Mitic et al., 2013; Leuner et al., 298 2004; Lifschytz et al., 2006; Monleón et al., 2002; Hodes et al., 2010). One possible explanation for our 299 lack of sex-specific effects stems from our general lack of sex differences in the behaviors analyzed, and 300 perhaps a baseline difference in behavior is necessary to elicit a sex-specific effect. The results of Mitic 301 et al. (2013); Leuner et al. (2004) and Lifschytz et al. (2006) in rats all observe sex-specific responses to 302 fluoxetine only when the sexes differed in behaviors without the drug. We do not have adequate data to 303 confirm this explanation and more experimentation along with physiological data would be necessary. 304

Another possible explanation for our lack of sex-specific drug effects could be our choice of dose. 305 Our choice of 1mg/L of nicotine is quite low compared with other studies in zebrafish (Levin et al., 2007; 306 Sackerman et al., 2010), and while our dosage of fluoxetine was much higher than is typically reported 307 (Egan et al., 2009; Wong et al., 2013), it is typically administered chronically. We would also like to note 308 that the sex-specific results of Faraday et al. (1999) utilizing nicotine in rats was only observed in one of 309 the two strains used. Zebrafish are highly genetically diverse (Parichy, 2015) and strain differences in 310 behavior (Benner et al., 2010; Egan et al., 2009) and drug efficacy (Sackerman et al., 2010) have been 311 reported. Therefore the possibility exists for sex-dependent drug effects to be observed in another strain. 312 Finally, we cannot dismiss the possibility that zebrafish simply don't exhibit sex-specific effects with 313 fluoxetine or nicotine. While there is no literature in this species to compare our results with, a recently 314 published study utilizing medaka (Oryzias latipes), another small teleost fish from southeast Asia, fails 315 to find sex-specific effects of chronic fluoxetine on many of the same behaviors described in the present 316 study (Ansai et al., 2016). More research is necessary to confirm any of the explanations given for our 317 lack of observed sex-drug interactions. The absence of studies considering sex-specific effects of drugs is 318 problematic if zebrafish are to remain a relevant model of pharmacology research. The topic has become 319 a concern in all animal models that NIH is going to start requiring all animal research to include sex as 320 part of the study unless deemed unnecessary (Clayton and Collins, 2014). If it turns out that strain is a 321 major factor influencing our results, then the abundance of genetically diverse populations could make 322 zebrafish an exciting tool to aid in the growing field of pharmakogenetics and personalized medicine in 323 which genetic background, among other traits, will be important for determining what drugs will be most 324 effective for treating disorders. 325

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329 **REFERENCES**

- Ansai, S., Hosokawa, H., Maegawa, S., and Kinoshita, M. (2016). Chronic fluoxetine treatment induces anxiolytic responses and altered social behaviors in medaka, Oryzias latipes. *Behav. Brain Res.*,
- 332 303:126–136.
- Barcellos, L. J. G., Ritter, F., Kreutz, L. C., Quevedo, R. M., da Silva, L. B., Bedin, A. C., Finco, J., and
- ³³⁴ Cericato, L. (2007). Whole-body cortisol increases after direct and visual contact with a predator in ³³⁵ zebrafish, Danio rerio. *Aquaculture*, 272(1-4):774–778.
- Barros, T. P., Alderton, W. K., Reynolds, H. M., Roach, a. G., and Berghmans, S. (2008). Zebrafish: an
- emerging technology for in vivo pharmacological assessment to identify potential safety liabilities in early drug discovery. *Br. J. Pharmacol.*, 154(7):1400–13.
- Beasley, C. M., Masica, D. N., and Potvin, J. H. (1992). Fluoxetine: a review of receptor and functional effects and their clinical implications. *Psychopharmacology* (*Berl*)., 107(1):1–10.
- Bencan, Z. and Levin, E. D. (2008). The role of alpha7 and alpha4beta2 nicotinic receptors in the
- nicotine-induced anxiolytic effect in zebrafish. *Physiol. Behav.*, 95(3):408–12.

- Bencan, Z., Sledge, D., and Levin, E. D. (2009). Buspirone, chlordiazepoxide and diazepam effects in a
 zebrafish model of anxiety. *Pharmacol. Biochem. Behav.*, 94(1):75–80.
- Benner, M. J., Drew, R. E., Hardy, R. W., and Robison, B. D. (2010). Zebrafish (Danio rerio) vary by
- strain and sex in their behavioral and transcriptional responses to selenium supplementation. *Comp. Biochem. Physiol. A. Mol. Integr. Physiol.*, 157(4):310–318.
- Benowitz, N., Hukkanen, J., and Jacob, P. (2009). Nicotine chemistry, metabolism, kinetics and biomarkers. *Handb. Exp. Pharmacol.*, 192(192):1–29.
- Buske, C. and Gerlai, R. (2014). Diving deeper into Zebrafish development of social behavior: analyzing high resolution data. *J. Neurosci. Methods*, 234:66–72.
- ³⁵² Cachat, J., Stewart, A., Grossman, L., Gaikwad, S., Kadri, F., Chung, K. M., Wu, N., Wong, K., Roy,
- S., Suciu, C., Goodspeed, J., Elegante, M., Bartels, B., Elkhayat, S., Tien, D., Tan, J., Denmark,
- A., Gilder, T., Kyzar, E., Dileo, J., Frank, K., Chang, K., Utterback, E., Hart, P., and Kalueff, A. V.
- (2010). Measuring behavioral and endocrine responses to novelty stress in adult zebrafish. *Nat. Protoc.*,
- 5(11):1786-1799.
- ³⁵⁷ Clayton, J. A. and Collins, F. S. (2014). Policy: NIH to Balance Sex in Cell and Animal Studies. *Nature*,
 ⁵⁰⁹(7500):282–283.
- Cohen, A., Young, R. W., Velazquez, M. a., Groysman, M., Noorbehesht, K., Ben-Shahar, O. M., and
- Ettenberg, A. (2009). Anxiolytic effects of nicotine in a rodent test of approach-avoidance conflict. *Psychopharmacology (Berl).*, 204(3):541–9.
- de Abreu, M., Koakoski, G., and Ferreira, D. (2014). Diazepam and Fluoxetine Decrease the Stress
 Response in Zebrafish. *PLoS One*, 9(7):1–5.
- Egan, R. J., Bergner, C. L., Hart, P. C., Cachat, J. M., Canavello, P. R., Elegante, M. F., Elkhayat, S. I.,
- Bartels, B. K., Tien, A. K., Tien, D. H., Mohnot, S., Beeson, E., Glasgow, E., Amri, H., Zukowska,
- Z., and Kalueff, A. V. (2009). Understanding behavioral and physiological phenotypes of stress and
- anxiety in zebrafish. *Behav. Brain Res.*, 205(1):38–44.
- Faraday, M. M., O'Donoghue, V. a., and Grunberg, N. E. (1999). Effects of nicotine and stress on startle amplitude and sensory gating depend on rat strain and sex. *Pharmacol. Biochem. Behav.*, 62(2):273–284.
- Gerlai, R. (2014). Social behavior of zebrafish: from synthetic images to biological mechanisms of shoaling. *J. Neurosci. Methods*, 234:59–65.
- Gerlai, R. (2015). Zebrafish phenomics: behavioral screens and phenotyping of mutagenized fish. *Curr. Opin. Behav. Sci.*, 2:21–27.
- Gerlai, R., Fernandes, Y., and Pereira, T. (2009). Zebrafish (Danio rerio) responds to the animated
- image of a predator: Towards the development of an automated aversive task. *Behav. Brain Res.*,
 201(2):318–324.
- Gilbert, D. (1979). Paradoxical Tranquilizing and Emotion-Reducing Effects of Nicotine. *Psychol. Bull.*,
 86(4):643–661.
- Graeff, F. G., Viana, M. B., and Mora, P. O. (1997). Dual role of 5-HT in defense and anxiety. *Neurosci. Biobehav. Rev.*, 21(6):791–799.
- Griffin, L. and Mellon, S. (1999). Selective serotonin reuptake inhibitors directly alter activity of neurosteroidogenic enzymes. *Proc. Natl. Acad. Sci.*, 96(23):13512–13517.
- Harris, S., Ramnarine, I. W., Smith, H. G., and Pettersson, L. B. (2010). Picking personalities apart:
- Estimating the influence of predation, sex and body size on boldness in the guppy Poecilia reticulata. *Oikos*, 119(11):1711–1718.
- Hodes, G. E., Hill-Smith, T. E., Suckow, R. F., Cooper, T. B., and Lucki, I. (2010). Sex-specific effects
- ³⁸⁸ of chronic fluoxetine treatment on neuroplasticity and pharmacokinetics in mice. *J. Pharmacol. Exp.* ³⁸⁹ *Ther.*, 332(1):266–273.
- King, A. J., Fürtbauer, I., Mamuneas, D., James, C., and Manica, A. (2013). Sex-differences and temporal
 consistency in stickleback fish boldness. *PLoS One*, 8(12):31–36.
- ³⁹² Kornstein, S. G., Schatzberg, a. F., Thase, M. E., Yonkers, K. a., McCullough, J. P., Keitner, G. I.,
- Gelenberg, a. J., Davis, S. M., Harrison, W. M., and Keller, M. B. (2000). Gender differences in
- treatment response to sertraline versus imipramine in chronic depression. *Am. J. Psychiatry*, 157:1445–
- зэ5 1452.
- Langheinrich, U. (2003). Zebrafish: a new model on the pharmaceutical catwalk. *Bioessays*, 25(9):904–12.
- Lesch, K. P., Bengel, D., Heils, A., Sabol, S. Z., Greenberg, B. D., Petri, S., Benjamin, J., Müller, C. R.,

- Hamer, D. H., and Murphy, D. L. (1996). Association of anxiety-related traits with a polymorphism in 398 the serotonin transporter gene regulatory region. Science (80-.)., 274(5292):1527-1531. 399
- Leuner, B., Mendolia-Loffredo, S., and Shors, T. J. (2004). Males and females respond differently to 400 controllability and antidepressant treatment. Biol. Psychiatry, 56(12):964–970. 401
- Levin, E. D., Bencan, Z., and Cerutti, D. T. (2007). Anxiolytic effects of nicotine in zebrafish. Physiol. 402 Behav., 90(1):54-8. 403
- Lifschytz, T., Shalom, G., Lerer, B., and Newman, M. E. (2006). Sex-dependent effects of fluoxetine and 404 triiodothyronine in the forced swim test in rats. Eur. Neuropsychopharmacol., 16(2):115–21. 405
- Martényi, F., Dossenbach, M., Mraz, K., and Metcalfe, S. (2001). Gender differences in the efficacy 406 of fluoxetine and maprotiline in depressed patients: A double-blind trial of antidepressants with 407 serotonergic or norepinephrinergic reuptake inhibition profile. Eur. Neuropsychopharmacol., 11:227– 408
- 232. 409
- Mathur, P. and Guo, S. (2010). Use of zebrafish as a model to understand mechanisms of addiction and 410 complex neurobehavioral phenotypes. Neurobiol. Dis., 40(1):66-72. 411
- Matta, S. G., Balfour, D. J., Benowitz, N. L., Boyd, R. T., Buccafusco, J. J., Caggiula, A. R., Craig, C. R., 412
- Collins, A. C., Damaj, M. I., Donny, E. C., Gardiner, P. S., Grady, S. R., Heberlein, U., Leonard, S. S., 413
- Levin, E. D., Lukas, R. J., Markou, A., Marks, M. J., McCallum, S. E., Parameswaran, N., Perkins, 414
- K. a., Picciotto, M. R., Quik, M., Rose, J. E., Rothenfluh, A., Schafer, W. R., Stolerman, I. P., Tyndale, 415
- R. F., Wehner, J. M., and Zirger, J. M. (2007). Guidelines on nicotine dose selection for in vivo research. 416
- Psychopharmacology (Berl)., 190(3):269–319. 417
- Maximino, C., de Brito, T. M., Colmanetti, R., Pontes, A. A. A., de Castro, H. M., de Lacerda, R. I. T., 418
- Morato, S., and Gouveia, A. (2010). Parametric analyses of anxiety in zebrafish scototaxis. *Behav.* 419 Brain Res., 210(1):1–7. 420
- Mehta, N. S., Wang, L., and Redei, E. E. (2013). Sex differences in depressive, anxious behaviors and 421 hippocampal transcript levels in a genetic rat model. Genes, Brain Behav., 12:695-704. 422
- Mitic, M., Simic, I., Djordjevic, J., Radojcic, M. B., and Adzic, M. (2013). Gender-specific effects of 423
- fluoxetine on hippocampal glucocorticoid receptor phosphorylation and behavior in chronically stressed 424
- rats. *Neuropharmacology*, 70:100–111. 425
- Monleón, S., Urquiza, A., Carmen Arenas, M., Vinader-Caerols, C., and Parra, A. (2002). Chronic 426
- administration of fluoxetine impairs inhibitory avoidance in male but not female mice. Behav. Brain 427 Res., 136(2):483-488. 428
- Oswald, M. E., Drew, R. E., Racine, M., Murdoch, G. K., and Robison, B. D. (2013a). Is Behavioral 429 Variation along the Bold-Shy Continuum Associated with Variation in the Stress Axis in Zebrafish? 430
- Physiol. Biochem. Zool., 85(6):718-28. 431
- Oswald, M. E., Singer, M., and Robison, B. D. (2013b). The quantitative genetic architecture of the 432 bold-shy continuum in zebrafish, Danio rerio. PLoS One, 8(7):e68828. 433
- Palanza, P. (2001). Animal models of anxiety and depression: how are females different? Neurosci. 434 Biobehav. Rev., 25(3):219-233. 435
- Parichy, D. M. (2015). Advancing biology through a deeper understanding of zebrafish ecology and 436 evolution. *Elife*, 4:1–11. 437
- Pavlidis, M., Digka, N., Theodoridi, A., Campo, A., Barsakis, K., Skouradakis, G., Samaras, A., and 438
- Tsalafouta, A. (2013). Husbandry of zebrafish, Danio rerio, and the cortisol stress response. Zebrafish, 439 10(4):524-31. 440
- Picciotto, M. R., Brunzell, D. H., and Caldarone, B. J. (2002). Effect of nicotine and nicotinic receptors 441 on anxiety and depression. *Neuroreport*, 13(9):1097–1106. 442
- Rasmussen, B. B., Menp, J., Pelkonen, O., Loft, S., Poulsen, H. E., Lykkesfeldt, J., and Brfsen, K. (1995). 443
- Selective serotonin reuptake inhibitors and theophylline metabolism in human liver microsomes: potent 444 inhibition by fluvoxamine. Br. J. Clin. Pharmacol., 39:151-159. 445
- Sackerman, J., Donegan, J. J., Cunningham, C. S., Nguyen, N. N., Lawless, K., Long, A., Benno, R. H., 446
- and Gould, G. G. (2010). Zebrafish Behavior in Novel Environments: Effects of Acute Exposure to 447
- Anxiolytic Compounds and Choice of Danio rerio Line. Int. J. Comp. Psychol., 23(1):43-61. 448
- Shin, J. T. and Fishman, M. C. (2002). From Zebrafish to human: modular medical models. Annu. Rev. 449 Genomics Hum. Genet., 3(24):311-40. 450
- Sison, M. and Gerlai, R. (2010). Associative learning in zebrafish (Danio rerio) in the plus maze. Behav. 451 452

- Wilson, A. D. M. and Godin, J.-G. J. (2009). Boldness and behavioral syndromes in the bluegill sunfish,
 Lepomis macrochirus. *Behav. Ecol.*, 20(2):231–237.
- ⁴⁵⁵ Wong, R. Y., Oxendine, S. E., and Godwin, J. (2013). Behavioral and neurogenomic transcriptome ⁴⁵⁶ changes in wild-derived zebrafish with fluoxetine treatment. *BMC Genomics*, 14(1):348.
- ⁴⁵⁷ Zhang, Y., Raap, D. K., Garcia, F., Serres, F., Ma, Q., Battaglia, G., and Van de Kar, L. D. (2000). Long-
- term fluoxetine produces behavioral anxiolytic effects without inhibiting neuroendocrine responses to
- conditioned stress in rats. *Brain Res.*, 855(1):58–66.
- Zimmerberg, B. and Farley, M. J. (1993). Sex differences in anxiety behavior in rats: Role of gonadal
- 461 hormones. *Physiol. Behav.*, 54(6):1119–1124.

462 FIGURES



Figure 1. Freezing behaviors (motionless at the bottom of the tank) appear not to be affected by exposure to fluoxetine or nicotine. These graphs show the probability of freezing \pm SE. (A) and the mean time spent frozen \pm SE (B) for both sexes in each drug treatment group. Females are represented as light bars and males as dark bars. The freezing probability was calculated from a logistic GLM and transformed back into probabilities for this figure using the 'lsmeans' package in R. Freezing time was transformed using a fourth root in order to meet the assumptions of normality in the ANOVA.



Figure 2. Average swimming speed (top) and consistency (individual variance) of swimming speed (bottom) are affected by fluoxetine and nicotine (A & B) as well as by sex (C & D). The fluoxetine treatment is not statistically different from the control, but is also not different from the nicotine treatment. Means \pm SE are reported. Results of the Tukey pairwise comparisons of drug treatment groups are delineated with letter groupings where similar letters represent a non-significant difference between treatments (p > 0.05). In panels A & B, females are represented with light bars and males with dark bars.



Figure 3. Average swimming depth (A) and average consistency (individual variance) of vertical usage (B) are affected by fluoxetine and nicotine. The nicotine treatment was not significantly different than the control with depth variance, but was also not different from the fluoxetine treatment. Means \pm SE are reported and the results of the Tukey pairwise comparisons of drug treatment groups are delineated with letter groupings where similar letters represent a non-significant difference between treatments (p > 0.05). Sex is distinguished by females with light bars and males with dark bars.



Figure 4. Average number of entries into the top half (A), latency to enter the top half (B), proportion of time spent in the top half (C), and proportion of time spent in center (D). Means \pm SE are reported and the results of the Tukey pairwise comparisons of drug treatment groups are delineated with letter groupings where similar letters represent a non-significant difference between treatments (p > 0.05). Sex is distinguished by females with light bars and males with dark bars. Latency to enter the top half is transformed using a fourth root transformation in order to meet the assumption of normality in the ANOVA.