Short- and Long-Term Functional Consequences of Fluoxetine Exposure During Adolescence in Male Rats

Sergio D. Iñiguez, Brandon L. Warren, and Carlos A. Bolaños-Guzmán

Background: Fluoxetine (FLX), a selective serotonin reuptake inhibitor, is prescribed for the treatment of major depressive disorder in young populations. Here, we explore the short- and long-term consequences of adolescent exposure to FLX on behavioral reactivity to emotion-eliciting stimuli.

Methods: Adolescent male rats received FLX (10 mg/kg) twice daily for 15 consecutive days (postnatal days 35–49). The influence of FLX on behavioral reactivity to rewarding and aversive stimuli was assessed 24 hours (short-term) or 3 weeks after FLX treatment (long-term). A separate group of adult rats was also treated with FLX (postnatal days 65–79) and responsiveness to forced swimming was assessed at identical time intervals as with the adolescents.

Results: Fluoxetine exposure during adolescence resulted in long-lasting decreases in behavioral reactivity to forced swimming stress and enhanced sensitivity to sucrose and to anxiety-eliciting situations in adulthood. The FLX-induced anxiety-like behavior was alleviated by re-exposure to FLX in adulthood. Fluoxetine treatment during adolescence also impaired sexual copulatory behaviors in adulthood. Fluoxetine-treated adult rats did not show changes in behavioral reactivity to forced swim stress as observed in those treated during adolescence and tested in adulthood.

Conclusions: Treating adolescent rats with FLX results in long-lived complex outputs regulated by the emotional valence of the stimulus, the environment in which it is experienced, and the brain circuitry likely being engaged by it. Our findings highlight the need for further research to improve our understanding of the alterations that psychotropic exposure may induce on the developing nervous system and the potential enduring effects resulting from such treatments.

Key Words: Adolescence, antidepressant, anxiety, depression, fluoxetine, rat, sexual behavior

Until relatively recently, the existence of major depressive disorder (MDD) in pediatric populations was not well recognized. Epidemiological reports now indicate that mood disorders are quite common early in life, affecting approximately 2% to 8% of children and adolescents, respectively (1,2). Pediatric MDD can lead to impairments in various psychiatric and functional domains such as antisocial personality, bipolar disorder, substance abuse, homelessness, self-harm, and up to 75% risk of recurrent depressive episodes in adulthood (3–7). These observations indicate an adverse impact of MDD on the development of neural substrates mediating cognitive, emotional, and social functioning (8–10). Thus, depression is a serious disorder necessitating timely and appropriate therapeutic intervention.

Fluoxetine (FLX) (Prozac), a selective serotonin reuptake inhibitor (SSRI), is the first drug approved for pediatric MDD (11). Although data about the effectiveness and safety of pharmacotherapy in youngsters are sparse, it is conceivable that treatment decisions for acute management of symptoms are necessitating timely and appropriate therapeutic intervention. Given the prevalence of prescription antidepressant use during adolescence and the scarcity of knowledge regarding long-term effects of such treatments, it is essential that the neurobiological consequences associated with FLX exposure be characterized. Thus, this study was designed to assess the short- and long-term behavioral responsivity to a range of emotion-eliciting stimuli after FLX exposure during adolescence (postnatal day PD 35–49) in male rats.

The acute effects of SSRI antidepressant medications are well defined: they increase the brain’s serotonin neurotransmission; however, they exert their mood-elevating effects after prolonged (i.e., weeks) administration (22,23). Serotonin is pivotal in the regulation of adolescent brain development in both rodents and humans (24,25). There is extensive serotonergic innervation of key brain regions involved in the control of emotional, cognitive, and motivated behaviors (25–28), and dysregulation of this neurotransmitter system has been correlated with deficits in behavior and emotional regulation (29–32). Because SSRI exposure in youngsters occurs at a time of ongoing neuronal adaptations (33–35) and such treatments can last for years (7,36,37), it is not difficult to conceive the notion that antidepressant treatments impact development of brain pathways dramatically influencing neurobiological functioning later in life.

Methods and Materials

Subjects
Male Sprague-Dawley rats were obtained from Charles River (Raleigh, North Carolina). For the initial experiment (Figure 1), rats arrived on the same day at PD30 (adolescent) and PD60 (~250–275 g, adults). For all other experimental conditions, rats arrived on PD30 and treatment started at PD35 or PD65 as depicted in Figure S1 in Supplement 1. The age at the start and duration of the experimental manipulations in adolescent rats (PD35–PD49) was selected because it roughly approximates adolescence in humans (33,35,38). Rats were housed in pairs in clear polypor
pylene boxes containing wood shavings in an animal colony maintained at 23°C to 25°C on a 12-hour light-dark cycle in which lights were on between 07:00 and 19:00 hours. Rats were provided with food and water ad libitum.

Drug Treatment and Experimental Design
Fluoxetine hydrochloride was obtained from Sigma (St. Louis, Missouri), dissolved in sterile distilled water, and administered in a volume of 2 mL/kg. An initial experiment was conducted using the forced swim test (FST) to establish the FLX dose that would reliably decrease immobility as characterized in adult (250 –275 g) rats (39,40). The FST consists of two swimming sessions over 2 days. The PD35 and PD65 rats were exposed to the FST on day 1 and then received intraperitoneal injections of FLX (0, 2.5, 5, 10, or 20 mg/kg) 23 hours, 5 hours, and 1 hour before re-exposure to the FST (day 2). Based on the results from this experiment (Figure 1), separate groups of PD35 rats were treated with FLX (0 or 10 mg/kg) twice daily (4 hours apart) for 15 consecutive days. Rats were randomly assigned to treatment and behavioral conditions, and the schedule of behavioral testing was counterbalanced among all groups (Table S1 in Supplement 1). Because rodents metabolize FLX about 10 times faster than humans (41), this drug schedule was selected to approximate FLX levels observed clinically. Short-term behavioral testing began 24 hours after the last injection, whereas long-term assessments started when subjects reached adulthood (Figure S1A in Supplement 1). Rats assigned to receive FLX in adulthood (treatment starting at PD65, Figure S1B in Supplement 1) were used as positive control rats (matched for drug treatment and testing time) only for the FST. Rats treated with FLX during adolescence and re-exposed to FLX as adults were tested on a single behavioral paradigm (i.e., food approach in a novel environment; Figure S5 and Table S1 in Supplement 1). Behavioral observations and analyses were performed by observers with no knowledge of the treatment conditions of each rat. All experiments were conducted in compliance with the 1996 National Institutes of Health Guidelines for the Care and Use of Laboratory Animals and approved by Florida State University Animal Care and Use Committee.

Sucrose Preference
The sucrose preference test (Figure S4 in Supplement 1) consisted of a two-bottle choice paradigm, as described previously (42) (full details in Supplement 1).

Locomotor Activity
Spontaneous locomotor activity was indexed as distance traveled (cm) in an open-field (OF) apparatus for 30 minutes (see Figure S3A,B in Supplement 1 for details and results).

Novel Object Approach
This test was conducted over 2 days. Rats were introduced to the OF for 30 minutes (day 1). On day 2, rats were placed in a corner of the OF containing a single food pellet (familiar rat chow) placed on a circular white filter paper (12 cm)

Figure 1. Acute effects of fluoxetine on forced swimming behaviors in adolescent (PD35; n = 8–9/dose) and adult (PD70+; n = 8–10/dose) male rats (A–E). (A) Latency to become immobile, (B) total immobility, (C) swimming counts, (D) climbing counts, and (E) floating counts of rats tested on the forced swim test after three injections (same dose) of fluoxetine (0, 2.5, 5, 10, or 20 mg/kg) 1, 5, and 23 hours between swims. Data were analyzed using individual one-way analyses of variance (p < .05) between the age groups. *Significantly different from vehicle control rats within the same age group. PD, postnatal day.
positioned in the center of the apparatus. Latency to approach the food and begin feeding was scored. The test ended immediately after rats started feeding or if they failed to approach food after 5 minutes, at which time they were placed back in their home cage with normal access to food and water.

**Elevated Plus-Maze**

The time spent and number of entries into the open arms of an elevated plus-maze (EPM) were assessed over 5 minutes, as previously described (42) (Supplement 1).

**Forced Swim Test**

The FST was conducted as previously described (45). Latency to immobility, total immobility, and behavioral counts (i.e., swimming, climbing, and floating) were recorded (details in Supplement 1).

**Sexual Behavior**

The sexual behavior experiments were carried out as previously described (46) under red light conditions between 13:00 and 18:00 hours. Male rats were given a 5-minute acclimation period to the testing arena, and testing was initiated by the introduction of a receptive female rat to the arena. Testing sessions (at PD80 and PD90, respectively) lasted 90 minutes (Supplement 1).

**Statistical Analyses**

Assignment of subjects to the various testing conditions was random. Behavioral data were analyzed using one-way or mixed-design (between and within variables) repeated analyses of variance (ANOVA) followed by Fisher’s least significant difference post hoc test. When appropriate, additional Student t tests were used to determine statistical significance of pre-post comparisons. Data are expressed as the mean ± SEM. Statistical significance was defined as $p < .05$.

**Results**

**Establishing FST Behavioral Reactivity**

Fluoxetine increased latency to immobility in adolescents ($F(4, 39) = 5.43, p < .001$; Figure 1A, left panel; $n = 8–9/group$). Rats receiving 10 or 20 mg/kg FLX displayed longer latencies to immobility when compared with control rats ($p < .05$). Fluoxetine had a tendency toward decreasing total immobility ($p = .07$; Figure 1B) and dose-dependently increased swimming counts ($F(4, 39) = 3.77, p < .01$; Figure 1C), while having no effect on climbing or floating counts (Figure 1D,E).

Fluoxetine dose-dependently increased latency to immobility in adults ($F(4, 39) = 8.88, p < .001$; Figure 1A, right panel; $n = 8–10/group$). Rats receiving 5, 10, or 20 mg/kg FLX displayed longer latencies to immobility ($p < .05$) and decreased total immobility ($F(4, 39) = 3.12, p < .02$) compared with control rats ($p < .05$; Figure 1B). Fluoxetine increased swimming counts ($F(4, 39) = 2.72, p < .04$; Figure 1C), without affecting climbing or floating counts.

**Effects of FLX on Body Weight**

Based on the results above, 10 mg/kg FLX was selected to treat adolescent and adult rats for 15 days (twice daily). Figure S2 in Supplement 1 shows the effects of FLX on body weight gain in PD35 (n = 18/group) and PD65 (n = 7–8/group) rats. A mixed-design repeated measures ANOVA revealed that FLX significantly decreased weight gain across days [main effect: $F(14, 476) = 930.75, p < .0001$], drug [main effect: $F(1, 34) = 11.67, p < .002$; Figure S2A inset in Supplement 1], and as a function of day by drug [interaction: $F(14, 476) = 25.51, p < .0001$] in adolescent rats (Figure S2A in Supplement 1). Although body weight increased with age, the FLX-treated adolescent rats displayed lower weights than control rats ($p < .05$). Similarly, FLX reduced body weight in adult rats (Figure S2B in Supplement 1) as a function of injection day [$F(14, 182) = 14.93, p < .0001$], drug [$F(1, 13) = 25.11, p < .0001$; Figure S2B inset in Supplement 1], and day by drug [$F(14, 182) = 18.05, p < .0001$]. Fluoxetine-treated adult rats displayed lower weights than control rats ($p < .05$).

**Effects of Chronic FLX on Sucrose Preference**

Fluoxetine did not influence total fluid intake (water + sucrose; Figure 2B) 24 hours after treatment (n = 15/group; short-term). Conversely, there was a main effect of sucrose [$F(1, 24) = 4.71, p < .04$; Figure 2A], with FLX-treated rats preferring sucrose only at the .25% concentration ($p < .05$). A separate ANOVA revealed that FLX treatment during adolescence increased sucrose preference in adulthood (Figure 2C; long-term), without affecting total fluid intake (Figure 2D; n = 15/group). Sucrose preference varied by sucrose concentration [main effect: $F(4, 112) = 145.56, p < .05$] and drug [main effect: $F(1, 28) = 9.08, p < .05$]. Fluoxetine treatment increased sucrose preference only at the .125%, .25%, and .5% concentrations ($p < .05$, respectively; Figure 2C).

**Effects of FLX on Anxiety-Like Behaviors**

**Elevated Plus-Maze.** Fluoxetine induced anxiety-like behaviors 24 hours after the last injection (short-term; n = 8/group) and in adulthood (long-term; n = 8/group). Fluoxetine significantly decreased percent time spent [$F(1, 14) = 11.03, p < .005$; Figure 3A, left panel] and percent entries [$F(1, 14) = 9.63, p < .008$; Figure 3B, left panel] in the open arms of the EPM. Similarly, rats tested in adulthood spent significantly less percent time in the open arms [$F(1, 14) = 21.93, p < .0001$; Figure 3A, right panel, but did not differ in percent entries into the open arms of the EPM (Figure 3B, right panel).

**Novel Object Approach in a Familiar Environment.** There were significant differences in the latency to approach a novel object 24 hours after treatment [t(17) = −2.16, $p < .05$]. Fluoxetine-treated rats took significantly longer to approach the object than control rats (Figure 4A; n = 9–10/group). Additionally, once the FLX-treated rats first approached the object, they spent significantly more time exploring it (Figure 4B) than control rats [t(17) = −3.59, $p < .02$]. A somewhat similar behavioral pattern was observed in rats tested in adulthood: FLX-treated rats displayed longer latencies to approach (Figure 4C; n = 14–15/group; t(27) = −2.32, $p < .03$) but showed no differences in time spent exploring the object (Figure 4D).

**Latency to Feed in a Novel Environment.** Fluoxetine-treated rats had significantly longer latencies to approach food in a novel environment 24 hours after treatment [t(16) = −4.24, $p < .05$; n = 9/group; Figure 4E, short-term] or in adulthood [t(10) = −2.35, $p < .05$; n = 6/group; Figure 4E, long-term]. We also assessed whether FLX could reverse these effects in a separate group of adult rats pretreated with FLX during adolescence. Repeated [5 days; t(10) = −3.8, $p < .05$], but not acute (1 day), FLX (10 mg/kg) reversed the aberrant latency to approach food in these rats (Figure 4F; n = 6/group).

**Effects of FLX on the FST**

We used the FST to assess rats’ responsiveness to stress 24 hours after treatment (Figure 5A–C) or when they reached

www.sobp.org/journal
adulthood (Figure 5D–F). Fluoxetine-treated rats displayed longer latencies to immobility \( t(9) = -6.1, p < .05 \) and decreased total immobility \( t(9) = 3.01, p < .05 \) compared with control rats 24 hours after treatment (Figure 5A,B; \( n = 5–6/\) group). Fluoxetine induced higher swimming \( t(9) = -3.87, p < .05 \) and climbing counts \( t(9) = -2.67, p < .05 \), with lower floating counts \( t(9) = 9.16, p < .05 \) than control rats. Fluoxetine-treated rats during adolescence and tested in adulthood also displayed a behavioral profile similar to the short-term group (Figure 5D–F; \( n = 15/\) group): longer latencies to immobility \( t(28) = -2.39, p < .02 \); Figure 5D), lower total immobility \( t(28) = 3.40, p < .05 \); Figure 5E), higher swimming counts \( t(28) = -3.78, p < .001 \), higher climbing counts \( t(28) = -3.34, p < .05 \), and lower floating counts \( t(28) = 3.35, p < .002 \); Figure 5F).

A separate group of adult rats was tested on the FST after chronic FLX (matched drug treatment and testing schedule, as with the adolescent group above; Figure 6A–F) to determine whether these FLX-induced effects on the FST are specific to adolescent treatment. These adult FLX-treated rats showed a similar behavioral profile as the FLX-treated adolescents only when tested 24 hours after the last injection (Figure 6A–C; \( n = 7/\) group): longer latencies to immobility \( t(12) = -4.35, p < .001 \); Figure 6A), decreased total immobility \( t(12) = 3.48, p < .005 \); Figure 6B), higher swimming counts \( t(12) = -4.42, p < .001 \), higher climbing counts \( t(12) = -4.25, p < .001 \); Figure 6C), and lower floating counts \( t(12) = 6.06, p < .0001 \); Figure 6C). Fluoxetine had no effects when the long-term adult group was tested 21 days after treatment (Figure 6D–F, \( n = 7/\) group).

**Discussion**

Antidepressants are often prescribed to pediatric populations (21); yet, there is a scarcity of knowledge regarding the short-term and/or long-lasting neurobiological consequences of such treatments during early life (11). Thus, this study was designed to assess enduring behavioral outcomes in response to rewarding and aversive situations resulting from repeated FLX exposure during adolescence in male rats. This approach was taken because serotonin and compounds that regulate its function interact with mesolimbic reward systems, part of the circuitry

**Effects of Adolescent FLX Exposure on Sexual Behavior**

Fluoxetine-exposed rats exhibited deficits in sexual activity when assessed in two separate 90-minute sexual behavior sessions (PD80 and PD90, respectively; Figure 7A–C; \( n = 10/\) group). A repeated measures (sex session) ANOVA indicated that mount latency varied only as a function of drug \( F(1,18) = 7.38, p < .01 \); Figure 7A). Fluoxetine-pretreated rats displayed longer mount latency than control rats at PD80 (\( p < .05 \); Figure 7A, left panel) but not at PD90 (Figure 7A, right panel). Fluoxetine also influenced ejaculation latency between the groups \( F(1,18) = 28.31, p < .001 \), with FLX-exposed rats displaying longer times to reach the first ejaculation at PD80 (\( p < .05 \); Figure 7B, left panel) and PD90 (\( p < .05 \); Figure 7B, right panel). Ejaculation frequency was affected by FLX \( F(1,18) = 20.01, p < .0001 \); Figure 7C), with FLX-exposed rats showing lower ejaculation frequency than control rats at both PD80 (\( p < .05 \)) and PD90 (\( p < .05 \)) sessions.
controlling emotional and motivated behaviors (47–52). We report that exposure to FLX during PD35 to PD49 leads to decreased responsiveness to stressful situations, increased sensitivity to natural reward, and anxiety-eliciting situations, including deficits in sexual behavior, in adulthood.

Exposure to FLX during adolescence increased rats’ normal sensitivity to sucrose (a natural reward) in adulthood, while only inducing a minimal increase in preference (at the .25% concentration) in rats tested 24 hours after treatment. Because antidepressants reduce body weight and caloric intake in animals and humans (53–55), decreases in sucrose preference were expected. However, the lack of changes in overall liquid intake (sucrose/water) between the groups indicates that increases in preference are likely due to the ability of FLX to alter rats’ responsiveness to the rewarding effects of sucrose in adulthood. Therefore, it is possible that the young rats tested short-term did not respond robustly to sucrose because of the ability of FLX to decrease caloric intake and palatability of sweet solutions (56,57). To further explore reward sensitivity after FLX administration, time spent exploring a novel object in a familiar environment was measured (43,58,59). Fluoxetine-treated adolescent rats spent significantly longer exploring the object 24 hours after treatment, indicating that interacting with the novel object was rewarding (60). However, no changes in object exploration were observed long-term and it consequently failed to complement the sucrose preference findings. Brain reward pathways, such as the nucleus accumbens (NAc) and its dopaminergic input from the ventral tegmental area, mediate responses to natural rewards (52,61,62). Ingesting sweet solutions and exploring novel objects activate this circuit (52,63,64) and disruption of this neural projection decreases interest for sucrose and novelty (61,65–67). As in the present study, research assessing the effects of antidepressant treatment on reward-related behavior reveals a complex picture. Antidepressants can decrease (68,69), increase (70,71), or have no effects (72) on responding for rewarding brain stimulation.

Figure 3. Fluoxetine (10 mg/kg, b.i.d.) exposure during adolescence regulates anxiety-like behavior in the EPM (A and B). Short-term (n = 8/group): FLX significantly reduced time spent (A, left panel) and entries (B, left panel) into the open arms of the EPM 24 hours after the last FLX injection (p < .05). Long-term (n = 8/group): FLX also reduced time spent (A, right panel) in the open arms of the EPM, without influencing entries (B, right panel) compared with VEH-treated control rats. Data are presented as percent time spent and percent entries (mean ± SEM) into the open arms of the EPM. b.i.d., twice daily; EPM, elevated plus-maze; FLX, fluoxetine; VEH, vehicle.

Figure 4. Effects of fluoxetine (10 mg/kg, b.i.d.) exposure during adolescence on the latency to approach a novel object in a familiar environment (A–D) and the latency to feed in a novel environment (E and F). Short-term (n = 9–10/group): FLX-treated rats had significantly longer latencies to approach (A and spent significantly more time exploring (B) the novel object 24 hours after the last FLX injection. Long-term (n = 14–15/group): FLX-treated rats displayed significantly longer latencies to approach (C) but spent similar time exploring (D) the novel object compared with control rats. (E) FLX increased latency to feed in a novel environment at both short-term (n = 9/group) and long-term (n = 6/group) time points of behavioral assessment. (F) Acute exposure to FLX (10 mg/kg) did not decrease latency to feed (F, left panel; n = 6/group) in a separate group of adult rats pretreated with FLX during adolescence. *Significantly different compared with VEH-treated control rats (p < .05). b.i.d., twice daily; FLX, fluoxetine; VEH, vehicle.

www.sobp.org/journal
with equivocal results when assessing responding for natural rewards (56,73). Nevertheless, antidepressants do sensitize brain reward pathways (74–76): they increase the firing activity of ventral tegmental area dopamine neurons (77), increase dopamine neurotransmission in the striatum (78–80), and enhance cocaine and morphine reward (81,82). Therefore, it is conceivable that FLX exposure during adolescence enhances reward processes that are likely discernable only in adulthood; however, more detailed studies assessing this notion are needed.

Our findings further indicate that FLX enhances reactivity to anxiogenic stimuli as measured in the EPM 24 hours after treatment in adolescent rats. This anxiety-like response was long-lived because the FLX-treated adolescent rats tested in adulthood showed similar anxiety-like responding. We also used latency to approach a novel object in a familiar environment and latency to start feeding in a novel environment as additional indexes of anxiety-like behaviors. When exposed to novel environments, rats face a conflict between their motivation to explore the environment (novelty preference) and fear of potential negative consequences (83,84). Thus, longer latencies to approach a novel object or to start feeding have been interpreted as indicative of higher levels of anxiety (29). Similar to the EPM findings, FLX-exposed rats took longer to approach a novel object in a familiar environment and to start feeding in a novel environment at both short- and long-term testing time points. Because familiarity of environment increases novelty seeking and the FLX-treated rats had longer latencies to approach the novel object in a familiar environment, it is conceivable that FLX exposure during adolescence induces “trait” and not situational anxiety (83,85); however, an alternate explanation could be that they have increased caution and less impulsivity (86). These results are supported by reports indicating that administration of SSRIs early in life results in long-lasting anxiogenic phenotypes (29,32,87). We also show that chronic, but not acute, re-exposure

**Figure 5.** Effects of fluoxetine (10 mg/kg, b.i.d.) on behavioral responsivity to swim stress (A–F). Short-term (n = 5–6/group): FLX-treated rats displayed significantly longer latencies to immobility (A), lower total immobility (B), higher swimming and climbing counts and lower floating counts (C) when compared with VEH-treated control rats. Long-term (n = 15/group): FLX-treated rats displayed similar behavioral profile (D–F) as those tested in the short-term condition when compared with their VEH-treated control rats. *Significantly different from VEH-treated rats (p < .05). Data are presented as latencies to become immobile and total immobility (in seconds) and as cumulative 5-second intervals of swimming, climbing, and floating counts (mean ± SEM). b.i.d., twice daily; FLX, fluoxetine; FST, forced swim test; VEH, vehicle.

**Figure 6.** Effects of fluoxetine (10 mg/kg, b.i.d.) treatment in adult rats (matched control group) on behavioral responsivity to forced swim stress (A–F). Short-term (n = 7/group): FLX-treated rats displayed significantly longer latencies to immobility (A), lower total immobility (B), higher swimming and climbing counts and lower floating counts (C) when compared with VEH-treated control rats. Long-term (n = 7/group): no differences were observed in any of the measures assessed between the groups. *Significantly different from VEH-treated rats (p < .05). Data are presented as latencies to become immobile and total immobility (in seconds) and as cumulative 5-second intervals of swimming, climbing, and floating counts (mean ± SEM). b.i.d., twice daily; FLX, fluoxetine; FST, forced swim test; VEH, vehicle.
Figure 7. Effects of fluoxetine (10 mg/kg, b.i.d.) exposure during adolescence in adult male rat sexual behavior (A–C, n = 10/group). Rats were given two 90-minute sessions (at postnatal day 80 and 90, respectively) to copulate with a receptive female. FLX treatment during adolescence increased the latency to mount an estrous receptive female (A), latency to reach the first ejaculation (B), and the total number of ejaculations (C) compared with VEH-treated control rats in the first sex session (PD80). During the second sex session (PD90), FLX treatment during adolescence increased latency to ejaculate (B, right panel) and decreased ejaculation frequency (C, right panel) without affecting latency to mount (A, right panel). *Significantly different from VEH-treated rats (p < .05). b.i.d., twice daily; FLX, fluoxetine; PD, postnatal day; VEH, vehicle.

(i.e., 5 days) to FLX in adulthood alleviates the FLX-induced anxiety-like behavior observed in the start-to-feeding test, findings consistent with previous reports (88). Furthermore, these findings are supported by studies showing that initial exposure to antidepressants, which have been used successfully for the management of anxiety disorders, exacerbate anxiogenic-like behaviors in humans (89–91) and animals (92–94), but these alterations dissipate after prolonged exposure (95–97). Under the appropriate conditions, behavioral reactivity in the OF can also be used as an index of anxiety (98); thus, it must be noted that the overall activity observed in the OF (Figure S3A,B in Supplement 1) does not complement our findings of increased anxiety-like behaviors.

Nevertheless, reports show that emotionality-related behavior from the OF and the EPM do not produce a common anxiety-related factor in adolescent rats (99), indicating that emotionality is multidimensional and that these tests do not always complement each other (100–103).

Fluoxetine-treated rats showed lower levels of behavioral despair when exposed to forced swimming. Rats tested 24 hours after treatment showed coping patterns commonly categorized as antidepressant-like behaviors (39,104,105), and this effect was also present in the long-term group (i.e., those treated during adolescence and tested in adulthood). These findings were not due to FLX-induced changes in motor activity because rats tested 24 hours after day 1 of FST showed no differences in distance traveled in the OF (Figure S3C,D in Supplement 1). An antidepressant-like phenotype after adolescent FLX counters reports showing that early-life (PD4–PD21) FLX administration renders mice vulnerable to stressful situations in adulthood (29,32,56). However, other studies using similar age and treatment regimen in mice also find equivocal results (87,88,106–108). To determine if these effects were specific to age of FLX exposure, we treated adult rats and exposed them to forced swimming 24 hours or 21 days after the last injection (i.e., matched drug regimen and testing time as the adolescents). Only those adult rats tested 24 hours after treatment displayed reduced behavioral despair in the FST, while the long-term group did not differ from control rats. Our results suggest that the FLX-induced effects in the FST may be specific to adolescent FLX treatment, and this assumption is supported by studies demonstrating that altered behavioral profiles induced by antidepressants are dependent on age of exposure (29,32,56,88). The mechanism(s) underlying these effects are unknown. In adults, antidepressants regulate complex cellular and intracellular signaling mechanisms such as brain-derived neurotrophic factor, extracellular signal-regulated kinase, and cyclic adenosine monophosphate-responsive element binding protein activity, factors associated with the regulation of mood and motivation, resulting in lasting synaptic changes influencing behavioral functioning (109–112). Fluoxetine actions in the nervous system are complex, and more detailed assessments of these phenomena accounting for length of exposure and discontinuation and developmental periods are clearly needed (35,97,113–115).

Lastly, we assessed whether FLX exposure during adolescence influences sexual behavior later in adulthood (see Figure S6 in Supplement 1). Fluoxetine-exposed rats showed increased latencies to mount and ejaculate and deficits in ejaculation frequency. Antidepressant treatments interfere with sexual functioning in both humans and rodents (116–118); however, these findings were unexpected, as the drug washout period for this particular group of animals was over 30 days and the behavioral deficits were observed at both PD80 and PD90 sessions. The mechanism(s) underlying these effects are also unknown. Serotonin interacts in a complex manner with several of its receptors to inhibit various aspects of sexual and ejaculatory functioning (119,120). Therefore, it is conceivable that early-life FLX induces long-lasting changes in receptors (e.g., increased sensitivity and/or density) known to inhibit sexual behavior (121). Alternatively, it is possible that sustained FLX exposure dysregulates second messenger systems, since others have shown that altered cyclic adenosine monophosphate-responsive element binding protein activity within the NAc of adult rats leads to impairments in the initiation of sexual behavior, but not the rewarding aspects of sex, in addition to increases in anxiety-like behavior (46,122,123). These findings parallel our results after adolescent FLX exposure: longer latencies to initiate sexual activity and...
increased sensitivity to anxiety-inducing situations in adulthood. Unfortunately, our results cannot discern whether the appetitive aspects of sexual behavior were influenced by FLX because the dependent variables assessed do not differentiate between inter- and effective treatment for pediatric MDD.

This work was supported by Grants R03DA020089 and R21DA022351 from the National Institute on Drug Abuse, a NARSAD Young Investigator Award, and a First Year Assistant Professor Award from Florida State University to CAB-G. SDI was supported by a McKnight Fellowship from the Florida Education Fund, a Neuroscience Fellowship from the Florida State University, and a National Research Service Award (F31DA027300) from the National Institute on Drug Abuse. BLW was supported by a Neuroscience Fellowship from the Florida State University. The authors report no biomedical financial interests or po-tential conflicts of interest.

Supplementary material cited in this article is available online.


www.sobp.org/journal


92. Bagdy G, Graf M, Anheuer ZE, Modes EA, Kantor S (2001): Anxiety-like effects induced by acute fluoxetine, sertraline or m-CPP treatment are reversed by pretreatment with the 5-HT2C receptor antagonist SB-242084 but not the 5-HT1A receptor antagonist WAY-100635. Int J Neurropsychopharmacol 4:399–408.


100. Lucki I (1997): The forced swimming test as a model for core and component behavioral effects of antidepressant drugs. Behav Pharmacol 8:523–532.


**Methods**

**Sucrose Preference**

The sucrose preference test consisted of a two-bottle choice paradigm (1). This paradigm has been used extensively to assess the effects of stress-induced anhedonia (2). Rats were habituated to drink water from two bottles for 5 days. At the start of the experiment, rats were exposed to ascending concentrations of sucrose (0, 0.125, 0.25, 0.5, and 1% wt/vol) for 2 days per sucrose concentration. Water and sucrose consumption were measured at 8:00 and 17:00 hours each testing day at which time the position of the sucrose bottle (left or right) was counterbalanced between the Fluoxetine- (FLX) and vehicle- (VEH) treated groups, across cages and days (see Figure S4, below). The preference for sucrose over water was used as a measure for rats’ sensitivity to reward.

**Locomotor Activity**

Spontaneous locomotor activity was assessed in an open-field (OF) apparatus that consisted of a square box (63 x 63 x 26 cm) that rats can explore freely. This apparatus is fully automated (Florida State University Psychology Department engineering group), and records the rats’ locomotor activity as ‘distance traveled’ in cm.

**Elevated Plus Maze**

FLX- and VEH- treated rats were tested for 5 min on the elevated plus maze (EPM), a behavioral model of anxiety-like behavior. The maze was made of gray plastic and consisted of two perpendicular, intersecting runways (12 cm wide X 100 cm long). One runway had tall walls (40 cm high) or “closed arms,” and the other one had no walls or “open arms.” The arms were connected together by a central area, and the maze was elevated 1 m from the floor. Testing was conducted between 9 AM and 1 PM under controlled light conditions (~90 lux). At the beginning of the 5-min observation, animals were placed in the central area, facing one of the open arms, and the cumulative time spent and number of entries into the open arms was recorded (3).
Forced Swim Test

The forced swim test (FST) is a 2-day procedure in which rats are forced to swim under conditions in which they cannot escape. On the first day, rats are forced to swim. Initially, they engage in escape-like behaviors but eventually adopt a posture of immobility in which they make only the movements necessary to maintain their head above water. When retested 24 h later, rats become immobile very quickly; however, antidepressant treatment between the forced swim exposures can significantly increase their escape-like behaviors, an effect that has been correlated with antidepressant activity in humans (4). At the start of the experiment, rats were placed in plastic cylinders (75 x 30 cm) filled to 54 cm depth with 25°C water and forced to swim for 15-min. At the end of this period, rats were removed from the water, dried with towels, and placed in a warmed enclosure for 30-min. All cylinders were emptied and cleaned between rats. Twenty-four h after the forced swim, rats were retested for 5-min under identical conditions, and sessions were videotaped. In this study, the latency (sec) to become immobile, total immobility (sec), and behavioral counts (floating, swimming, and climbing) were the dependent variables [see (5,6)]. Behavioral counts were rated at 5-sec intervals during the 5-min retest. Latency to immobility was defined as the time at which the rat first initiated a stationary posture that did not reflect attempts to escape from the water (7). To qualify as immobility, this posture had to be clearly visible and maintained for ≥2.0 sec.

Sexual Behavior

The sexual behavior experiments were carried out as previously reported (8-10). Rats were housed in a separate room maintained on a 12-h light/dark cycle (lights on between 24:00 and 12:00 h). Sexual behavior was assessed under red light conditions between 13:00-18:00 h in a circular arena (60 cm) containing wood chips on the floor. Each male was given a 5 min acclimation period to the testing arena. Testing started at the end of the acclimation period by the introduction of a receptive female to the arena. Testing sessions (at PD80 and PD90, respectively) lasted 90-min (see Figure S6 below). Behaviors recorded were mount latency (ML), elapsed time between introduction of the female and the first display of mounting, ejaculation latency (EL), time between first mount and first ejaculation, ejaculation frequency (EF), and total number of ejaculations. For rats that either did not display mounting behavior or failed to reach an ejaculation during the test session, the ML and EL was given as 90 min [similarly to (9)]. Sprague-Dawley ovariectomized female rats (Charles River, Raleigh, NC) were used in these experiments. Receptivity of the females was induced by injection of estradiol benzoate [50
mg, subcutaneously (sc) and progesterone (500 mg, sc) 48 and 4-6 h before testing, respectively. One week prior to the experiment, the females were tested for one intercourse session with an experienced male. Prior to testing, female receptivity was verified by the exhibition of lordosis, in the presence of the experienced male, and accepted intromission. Each female was used to test only one experimental male.

**Results**

*Effects of FLX on Basal Locomotor Activity*

Chronic VEH- or FLX (10 mg/kg, b.i.d.) exposure during adolescence did not influence distance traveled (cm) in the open field 24 h after treatment (Short-term; Figure S3-A; n=10/group), or in adulthood (Long-term; Figure S3-B; n=14-15/group).

*Basal Locomotor Activity 24 h After Day 1 of FST*

Because changes in FST performance can be influenced by differences in motor activity, separate groups of VEH- and FLX-treated rats were tested in the OF 24 h after day 1 of FST (n=6/group). No changes in locomotor activity were evident in rats tested either short- or long-term (Figure S3 C-D) after treatment.
Table S1. Experimental/Treatment Design for Adolescent rats treated with Fluoxetine. Adolescent rats received fluoxetine (FLX; 10 mg/kg) or vehicle (VEH) intraperitoneal injections [twice daily (b.i.d.)] from PD35-49. Rats were tested in no more than 2 behavioral paradigms in the order of testing and time interval between tests as depicted in the Table. Rats were tested either 24 h (Short-term; PD50) or 20+ days after the last injection (Long-term; PD70+). Rats assigned to sex behavior were tested at PD80 and PD90, respectively. Rats in groups 8 and 9 were re-exposed to FLX as adults (>PD60): those in the acute condition (group 8) received a single injection of FLX or VEH at PD69, whereas those in the chronic condition (group 9) received a single injection of FLX or VEH for 5 consecutive days (PD65-69; see also Figure S5). Latency to start feeding behavior testing started at PD70.

<table>
<thead>
<tr>
<th>Group</th>
<th>Day 35-49</th>
<th>Interval</th>
<th>TEST 1</th>
<th>Interval</th>
<th>TEST 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>VEH or FLX, b.i.d.</td>
<td>24 hours</td>
<td>FST</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>2</td>
<td>VEH or FLX, b.i.d.</td>
<td>24 hours</td>
<td>Sucrose Preference</td>
<td>20 days</td>
<td>FST</td>
</tr>
<tr>
<td>3</td>
<td>VEH or FLX, b.i.d.</td>
<td>24 hours</td>
<td>EPM</td>
<td>20 days</td>
<td>Sucrose Preference</td>
</tr>
<tr>
<td>4</td>
<td>VEH or FLX, b.i.d.</td>
<td>24 hours</td>
<td>Latency to Start Feeding</td>
<td>20 days</td>
<td>EPM</td>
</tr>
<tr>
<td>5</td>
<td>VEH or FLX, b.i.d.</td>
<td>24 hours</td>
<td>Novel Object Approach</td>
<td>20 days</td>
<td>Latency to Start Feeding</td>
</tr>
<tr>
<td>6</td>
<td>VEH or FLX, b.i.d.</td>
<td>20 days</td>
<td>Locomotor Activity</td>
<td>24 hours</td>
<td>Novel Object Approach</td>
</tr>
<tr>
<td>7</td>
<td>VEH or FLX, b.i.d.</td>
<td>24 hours</td>
<td>Locomotor Activity</td>
<td>30 and 40 days</td>
<td>Sex Behavior</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Group</th>
<th>Day 35-49</th>
<th>Interval</th>
<th>Fluoxetine Re-exposure</th>
<th>Interval</th>
<th>TEST PD70</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>VEH or FLX, b.i.d.</td>
<td>20 days</td>
<td>Acute: PD69</td>
<td>24 hrs</td>
<td>Latency to Start Feeding</td>
</tr>
<tr>
<td>9</td>
<td>VEH or FLX, b.i.d.</td>
<td>16 days</td>
<td>Chronic: PD65-69</td>
<td>24 hrs</td>
<td>Latency to Start Feeding</td>
</tr>
</tbody>
</table>

FST, forced swim test; EPM, elevated plus maze
Figure S1. Timeline of developmental experimental procedures. All rats arrived in the laboratory on postnatal day (PD) 30. Rats were randomly assigned to receive fluoxetine (FLX) from (A) PD35-49 (adolescence) or from (B) PD65-79 (adulthood). All rats received intraperitoneal injections of FLX (10 mg/kg) or VEH twice daily (b.i.d.) 4 hrs apart (900 and 1300 hrs, respectively). Behavioral testing of rats treated during adolescence (A) was conducted either 24 hr after the last injection (PD50; Short-term) or when they reached adulthood (PD70; Long-term). Rats receiving FLX in adulthood (B) were tested either 24 hr (PD80; Short-term) or 21 days (PD100; Long-term) after the last injection.
Figure S2. Effects of repeated (15 days) exposure to fluoxetine (FLX; 10 mg/kg, b.i.d.) on weight gain. Data were analyzed by mixed-design (within: day of injection, between: FLX) repeated measures ANOVA followed by post hoc test. (A) Adolescents (n=18/group): body weight increased across days regardless of condition, and FLX treatment resulted in significantly lower weight gain, starting on day 5 of drug exposure, when compared to control rats. (B) Adults (n=7-8/group): similar pattern of results was obtained from the adult rats treated with FLX, resulting in lower weight gain starting on day 3 of drug exposure as compared to controls. *Significantly different when compared to VEH-treated controls (p<0.05). Data are presented as average weight gain across days and drug treatment (mean ± SEM, in grams).
Figure S3. Exposure to FLX [10 mg/kg (twice daily; b.i.d.)] during adolescence did not affect total basal locomotor activity in the open field in rats tested either 24 hr (A: Short-term; n=10/group) after the last injection, or when tested in adulthood (B: Long-term; n=14-15/group). Similarly, FLX treatment did not affect locomotor activity 24 h after day 1 (C and D) of forced swimming (FST; n=6/group).
Iñiguez et al.

**Figure S4.** Timeline: Sucrose Preference testing. All rats arrived in the laboratory on postnatal day (PD) 30. Rats were randomly assigned to receive fluoxetine (10 mg/kg twice daily) or vehicle from PD35-49. Rats assigned to the Short-term testing condition (A) were habituated to drink water from two water bottles starting on PD45 for five consecutive days. Twenty-four hrs after the last injection (PD50), rats were introduced to ascending concentrations of sucrose (SUC; 0.125, 0.25, 0.5 and 1%; two days per concentration). Rats assigned to the Long-term condition (B) were acclimated to drink water from two different bottles starting at PD65. At PD70, these rats were introduced to the same ascending concentrations of sucrose.
Fluoxetine Re-exposure Timeline: Latency to Feed in a Novel Environment test. Rats arrived in the laboratory on postnatal day (PD) 30. Rats were randomly assigned to receive either fluoxetine [FLX; 10 mg/kg twice daily (b.i.d.) ; n=12] or vehicle (n=12) from PD35-49. After treatment, rats were randomly assigned to either an acute (A) or chronic (B) FLX re-exposure treatment as adults. Rats in the Acute FLX re-exposure group received a single FLX (10 mg/kg) or vehicle injection on PD69, whereas the rats in the chronic FLX re-exposure group received once daily injections of FLX (10 mg/kg) or vehicle for five consecutive days (PD65-69). All rats were then tested on the latency to start feeding in a novel environment test on PD70.
**Figure S6.** Timeline: Sexual Behavior testing. All rats arrived in the laboratory on postnatal day (PD) 30. Rats were randomly assigned to receive either fluoxetine (10 mg/kg twice daily; n=10) or vehicle (n=10) from PD35-49. All rats were tested on sexual copulatory behaviors at PD80 (Session 1) and again at PD90 (Session 2).


