INTRODUCTION

The precise role of central serotonergic system in acoustic startle response (ASR) and prepulse inhibition (PPI) is not clearly defined. 8-OH-DPAT, a 5-HT1A receptor agonist, has been used to investigate this topic in rat models as it was reported to increase the magnitude of the startle response and decrease the PPI. Sipes and Geyer later confirmed the central involvement of 5-HT1A receptor in gating function by demonstrating that the reduction in PPI produced by intra-raphe microinjections of 8-OH-DPAT was prevented by a systemic injection of a selective 5-HT1A antagonist, (+) WAY 100135. These findings suggest that rats with enhanced 5-HT1A activity are inferior to execute sensorimotor gating function when encountering an acoustic startle stimulus.

Biological or environmental interferences during the developmental process have been proved important in determining the integrity of central 5-HT system is important to 5-HT1A-modulated sensorimotor gating in isolation-reared rats.
ing the sensorimotor gating function. For example, a neonatal 5-HT\(^+\) depletion by pharmacological intervention with 5,7-DHT strengthens the ASR, but leaves the PPI unaffected.\(^7\) However in a non-pharmacological approach, rats reared in social isolation since weanling characteristically exhibit a higher ASR and a lower PPI.\(^6\) The isolation rearing (IR)-induced gating deficit is developmentally specific, as it is unable to be ameliorated by re-socialization after so called “critical period” of early life.\(^9\) IR paradigm is thus appropriate in exploring the mechanism underlying the 5-HT\(^+\)A modulation of sensorimotor gating function, in particular social interaction is reported to affect 5-HT\(^+\)A profiles.\(^10\)

The interactive effects between IR and 5-HT\(^+\)A receptors appear inconsistent in rodent study. IR has been reported to augment the behavioral effects of 8-OH-DPAT in rats, possibly via a mechanism of supersensitivity of the post-synaptic 5-HT\(^+\)A receptor.\(^11\) On the other hand, 5-HT\(^+\)A activation may reverse IR-induced PPI deficit without any effect on the ASR in mice, via a possible mechanism of presynaptic 5-HT\(^+\)A autoreceptors.\(^12\) It is worth investigating whether 5-HT\(^+\) depletion model, as it downgrades the presynaptic 5-HT\(^+\) effects, could be used to explore the role of 5-HT\(^+\)A receptor in the IR-impaired gating function.

The present study examined the effects of 5HT\(^+\)A activation on ASR and PPI in isolation reared rats following 5-HT depletion by 5,7-DHT. The results may be helpful for a better understanding in terms of the functional role of 5-HT\(^+\)A receptor in developmentally social attachment.

**METHODS**

**Animals**

Male Sprague-Dawley (SD) rats aged at 21–23 days old (weaned; BioLASCO Taiwan Co., Ltd., Yilan, Taiwan) were used in the study and were divided randomly into 2 rearing conditions [social rearing (SOC, 2 rats per cage) vs. isolation rearing (IR, housed singly)]. The animals were kept in a temperature\(--(22 \pm 4^\circ\)C\) and humidity–controlled (50\(\pm\)20\%) room under a 12-h light/dark cycle (lights on from 07:00 to 19:00) and given ad libitum access to a standard laboratory chow diet (Ralston Purina, St. Louis, MO, USA) and sterile water. At their age of postnatal days 77–79,\(^13\) rats in each rearing condition were randomly assigned to one of the two managements (sham control vs. 5-HT depletion surgery), thus made it totally 4 groups (SOC-sham, IR-sham, SOC-depletion, IR-depletion, N=8 for each) to enter the behavioral protocol (7 days after the surgery) for measuring their ASR and PPI under different doses of 8-OH-DPAT. Locomotor activity was also monitored in both SOC and IR rats under 5-HT depleted condition. At the end of study, animals were sacrificed for analyzing their tissue levels of 5-HT. The study was approved by the Institutional Animal Care and Use Committee of National Defense Medical Center with the certificate number NDMC-10319.

**Locomotor activity**

Total travel distance was used to reflect the locomotor activity, and was measured using a computerized automated activity monitoring system (MED Associates, Inc., St. Albans, VT, USA). The system included four plexiglass chambers (43\(\times\)43\(\times\)30 cm) equipped with an infrared array of 16 photodetectors and corresponding light sources that emitted photo-beams 3 cm apart and 4.5 cm above the chamber floor. Travel distance was recorded at the assigned intervals and analyzed with Med-Associates software (MED Associates, Inc., St. Albans, VT, USA). Locomotor activity data were individually recorded every 5 min and expressed as cumulative 60-min data.\(^14\)

**PPI testing**

The PPI experiments were carried out in four startle chambers (SR-LAB, San Diego Instrument, San Diego, CA, USA), each with a plexiglass cylinder (9 cm diameter). A speaker mounted 24 cm above the cylinder provided the background noise, prepulse stimuli and startle stimuli, which were controlled with the SR-LAB program. Startle responses were transduced by a piezoelectric accelerometer mounted below the cylinder, digitized and rectified. The movement of the rats was measured for 100 ms after startle stimulus onset. The rats were allowed to habituate to the background noise of 65 dB for 5 min after being placed into the chambers. The acoustic stimuli were calibrated every week using a digital sound level meter (RadioShack, Fort Worth, TX, USA) that was positioned inside the cylinder on the startle platform. Sixty trials were conducted in each test session, with an average inter-trial interval of 15 s. The first and last 6 trials (in Block 1 and Block 3, respectively) each consisted of a single 40 ms 120 dB startle stimulus (i.e., startle stimulus alone). The middle 48 trials (Block 2) consisted of random delivery of 12 trials of startle stimulus alone, 12 no-stimulus trials, and 24 prepulse trials (i.e., prepulse stimulus with startle stimulus). The prepulse trials consisted of a single 120 dB startle stimulus preceded by a 20 ms non-startling prepulse stimulus of 12 dB above the background noise. The prepulse stimulus preceded the startle stimulus by 100 ms. The startle amplitude (measured in arbitrary units) of the first 6 trials (i.e., startle stimulus alone trials). The PPI was determined with the data from Block 2 according to the following formula: [1-{(startle amplitude following prepulse+pulse pair/startle amplitude following startle stimulus alone)}\()\times\)100\%.\(^15\)
5-HT tissue levels in cortex, striatum, and hippocampus

Rats were sacrificed by decapitation and their brains were rapidly removed. Various brain areas, including cortex, striatum, and hippocampus were rapidly dissected on an icy cold plate, then weighed and stored at -80°C until homogenization with ultrasonication in 0.2 mL of 7N perchloric acid (Sigma Chemical Industries, Ltd., Saint Louis, MO, USA). Homogenates were centrifuged at 12,000×g for 30 min at 4°C. The concentration of serotonin (5-HT) was determined by high performance liquid chromatography (HPLC) equipped with electrochemical detector (ECD, LC-4C, BAS, West Lafayette, IN, USA) (10 nA, filter 2.0 Hz, AppE cell 0.750 V), autosampler (Shimadzu SIL-10ADvp autosampler, Shimadzu, Japan). The supernatant was filtered through a 0.22 um filter, and was analyzed by HPLC equipped with an Alltima™ (Grace Davison Discovery Sciences, Ltd., Deerfield, IL, USA) reversed-phase C18 column (4.6×150 mm, 5 μm). Injection volume was 20 μL. The mobile phase contains 100 mM NaH2PO4.H2O, 0.74 mM sodium octanesulfonate (SOS), 0.02 mM EDTA and 20% methanol adjusted the pH to 3.0 by H3PO4 and with a flow rate of 1.0 mL/min. The calibration curve was obtained based on 6 levels (20, 40, 60, 80, 100, and 200 ppm) of 5-HT standard (Sigma Chemical Industries, Ltd.). The coefficient of determination (r²) was greater than 0.995.

Surgery

Microinjection needle (Hamilton, 30-gauge, 10 µL) was used to inject 5,7-DHT into ICV (AP: -0.8 mm from bregma, L: ±1.5 mm from the midline, DV: -0.4 mm from dura) with a flow rate of 1 µL/min controlled by a microinjection pump (CMA/100, Carnegie, Medicin, Sweden) for 10 min.

Drugs

Serotonergic neurotoxin 5,7-DHT (200 µg, Sigma-Aldrich, St. Louis, MO, USA) was dissolved in saline solution with 0.6% ascorbic acid and was administered via intracerebroventricular (ICV) injection. 30 min before the 5,7-DHT injection, desipramine (25 mg/kg, Sigma-Aldrich), a reuptake inhibitor of norepinephrine, was administered intraperitoneally to protect noradrenergic neurons from neurotoxicity. To activate the 5-HT1A receptors, 8-OH-DPAT (Sigma-Aldrich) was intraperitoneally injected (0, 10, and 100 µg/kg, in a volume of 1 mL/kg) 10 min prior to the behavioral testing.

Data analyses

Statistical analyses were performed across the groups via a multi-factor analysis of variance (ANOVA) by using SPSS Version 18.0 (SPSS Inc., Chicago, IL, USA), with REARING CONDITION and LESION as between-subject factors. Further analyses with post-hoc multiple comparisons were performed where possible. All data were expressed as mean±standard error of the mean (SEM). A p value of <0.05 was considered statistically significant.

RESULTS

Effects of 5,7-DHT on tissue 5-HT levels

The HPLC method confirmed the reductions of tissue 5-HT caused by 5,7-DHT in cortex, striatum, and hippocampus in which IR rats were less affected (for SOC rats, cortex: 32%, striatum: 35%, hippocampus: 33%; for IR rats, cortex: 73%, striatum: 51%, hippocampus: 61%) (Table 1). There were main effects of LESION [F(1,16)=9.61, p<0.01] and REARING CONDITION [F(1,16)=5.06, p<0.05] in striatum, and main effects of LESION [F(1,16)=9.40, p<0.01] and REARING CONDITION [F(1,16)=5.24, p<0.05] in hippocampus with REARING CONDITION×LESION interaction [F(1,16)=4.88, p<0.05]. For cortex, there was a main effect of LESION [F(1,16)=6.15, p<0.05].

Effects of 5,7-DHT on locomotor activity

IR rats exhibited greater locomotor activity (indexed by total travel distance) in the 60 min activity test [t(30)=3.4, p<0.01]. The treatment of 5,7-DHT did not cause any effect on locomotor activity in rats of both rearing conditions [no effects of LESION or REARING CONDITION×LESION; main effect of REARING CONDITION remained, F(1,28)=9.02, p<0.01] (Figure 1).

Effects of 5,7-DHT on ASR and PPI

Both REARING CONDITION and LESION have no effects on the ASR. For PPI, IR rats was inferior than SOC rats [main effects of REARING CONDITION, F(1,28)=7.98, Table 1. 5-HT concentration (pg/mg) in cortex, striatum, and hippocampus (N=5 for each group)
p<0.01]. The treatment of 5,7-DHT did not cause any effect on PPI in rats of both rearing conditions [no effects of LESION or REARING CONDITION×LESION] (Figure 2).

Effects of 8-OH-DPAT on ASR and PPI in 5-HT depleted rats reared in social isolation

For ASR, 8-OH-DPAT has no effect on the ASR in SOC rats. However, it strengthens the ASR in both IR-sham and IR-depletion rats [for IR rats, main effect of DOSE, F(2,28)=6.11, p<0.01] (Figure 3).

For PPI, LESION was a key factor determining the 8-OH-DPAT effect, as the drug at 100 µg/kg caused a higher PPI in 5-HT depleted SOC rats than their sham controls [for SOC rats, main effect of LESION, F(1,14)=4.90, p<0.05 and effect of LESION×DOSE F(2,28)=3.73, p<0.05]. However, this effect was reversed in an opposite manner for IR rats, in which the 8-OH-DPAT at 100 µg/kg strengthens the PPI in non-lesioned IR rats but downgrades the PPI in lesioned IR rats [for IR rats, main effect of lesion, F(1,14)=9.38, p<0.01 and effect of lesion×dose F(2,28)=5.52, p<0.01] (Figure 4).

**DISCUSSION**

The major finding of our study is that 5-HT depletion may lead to distinctive 8-OH-DPAT effects on PPI in rats, depending on their rearing conditions. In the present study, 8-OH-DPAT at 100 µg/kg improved PPI in 5-HT depleted SOC rats, whereas it downgraded PPI in the 5-HT depleted IR rats. In other words, the integrity of 5-HT neuron is important for the 5-HT1A modulatory effects on sensorimotor gating function, in which the development of social attachment may serve as a key factor in determining the effects.

Locomotor hyperactivity and impairment of PPI are two prominent characteristics in rodent model of IR, and were firstly validated in the present study. Rather than hyperactivi-

![Figure 1](image1.png)

**Figure 1.** Effects of 5-HT depletor, 5,7-DHT on locomotor activity in isolation reared rats after 5HT depletion. A: IR rats are hyperactive than social controls. B: 5,7-DHT has no effects on locomotor activity in rats of both rearing conditions. Data are represented by mean±SEM. *p<0.01, N=16 for SOC and IR rats (A) and N=8 for rats under different rearing and treatment conditions (B). 5-HT: serotonin, SOC: social, IR: isolation-rearing.

![Figure 2](image2.png)

**Figure 2.** Effects of 5-HT depletor, 5,7-DHT on ASR and PPI in isolation reared rats after 5HT depletion. A: Both REARING CONDITION and LESION have no effects on the ASR. B: IR rats perform lower PPI than social controls; 5,7-DHT has no effects on PPI in rats of both rearing conditions. Data are represented by mean±SEM. *p<0.01, N=8 for each group. 5-HT: serotonin, SOC: social, IR: isolation-rearing, ASR: acoustic startle response, PPI: prepulse inhibition.
ty, PPI is more appropriate in addressing developmental specificity because the IR-induced PPI deficit is unable to be reversed by resocialization later. The present study demonstrated that the depletion of central 5-HT did not modulate the IR effects on both locomotor activity and sensorimotor gating function (including ASR and PPI). Note the condition of 5-HT depletion leads to a damage to central 5-HT-containing axons and terminals, however the 5-HT efflux appeared unaffected in lesioned rats compared to the sham control rats, in both the baseline and the anxiety triggered conditions. A compensatory mechanism is therefore highly suggested for the maintenance of synaptic function. Our data implies that this mechanism is preserved in rats reared in social isolation.

Figure 3. Effects of different doses of 8-OH-DPAT on ASR in isolation reared rats after 5HT depletion. A: 8-OH-DPAT has no effect on the ASR in SOC rats (comparisons were made among upper, middle and bottom panels). B: 8-OH-DPAT strengthens the ASR in both IR-sham and IR-depletion rats (comparisons were made among upper, middle and bottom panels). Data are represented by mean±SEM, N=8 for each group. 5-HT: serotonin, SOC: social, IR: isolation-rearing, ASR: acoustic startle response.

Figure 4. Effects of different doses of 8-OH-DPAT on PPI in isolation reared rats after 5HT depletion. A: LESION was a key factor determining the 8-OH-DPAT effect, as the drug at higher dose (i.e., 100 µg/kg) caused a higher PPI in 5-HT depleted SOC rats than their sham controls (comparisons were made among upper, middle and bottom panels). B: 8-OH-DPAT strengthens the PPI in non-lesioned IR rats. For lesioned IR rats, higher dose (100 µg/kg) 8-HDPAT downgrades the PPI (comparisons were made among upper, middle and bottom panels). Data are represented by mean±SEM. *p<0.01, N=8 for each group. 5-HT: serotonin, SOC: social, IR: isolation-rearing, PPI: prepulse inhibition.
The most striking finding of the present study is the effect of 8-OH-DPT on sensorimotor gating function in IR rats, particularly in the lesioned condition. For ASR, 8-OH-DPT boosted it in IR but not SOC rats. 8-OH-DPT was previously reported to increase the amplitude of startle in a similar acoustic paradigm. Since in the present study the 8-OH-DPT increased ASR displayed in both the sham control and the 5,7-DHT pretreated rats, the effect should be less relevant to the lesion condition of 5-HT neurons. In other words, the nature of hyper-reactivity to startle stimulus of IR rats is presumably more relevant to 5-HT1A activation than the integrity of 5-HT neurons.

In contrast to ASR, the IR-induced impairment of PPI represents a failure of information-filtering to mitigate the response to the ensuing stimuli (i.e., a dysfunction of the central inhibitory ability), and the integrity of 5-HT neurons rather than rearing condition became the key to affect the 8-OH-DPT effects on PPI. High dose of 8-OH-DPT (at 100 µg/kg) caused a greater PPI in 5-HT depleted SOC rats than their sham controls, indicating that 5-HT1A activation-downgraded PPI can be modulated if 5-HT neurons are not intact, possibly via a compensatory mechanism as mentioned above. However, the mechanism appears different in the IR rats, as in the present study the 8-OH-DPT strengthened the PPI in IR-sham rats but downgraded it in IR-lesioned rats. In other word, processes responsible for maintaining the function of synaptic 5-HT transmission of IR rats in the condition of 5,7-DHT impact have altered now due to the activation of 5-HT1A receptors.

The mechanisms underlying these observations might be complicated, however several points are worth addressing. First, it implies that the two pharmacological managements (5,7-DHT and 8-OH-DPT) exert their effects on sensorimotor gating in IR rats via different pathways. While 5,7-DHT exerts its effects presynaptically, 8-OH-DPT exerts its effects pre and post-synaptically, because 5-HT1A receptors may be expressed in both places. Second, 8-OH-DPT at the dose used in the present study (i.e., 100 µg/kg) is generally inadequate to activate the postsynaptic 5-HT1A receptors, but since the IR rats characteristically may have a supersensitive profile of the post-synaptic 5-HT1A receptor, the 5,7-DHT-readjusted presynaptic function may affect the tonicity/balance between pre and post-synaptic 5-HT1A receptors, and therefore responsible for the diverse effects of 8-OH-DPT on PPI in IR-sham and IR-depletion rats. Finally, although 8-OH-DPAT is a potent agonist at 5-HT1A receptors, it may also antagonize the effects of 5-HT itself, suggesting an effect of partial agonism, which is along with the dose-dependent, biphasic effects of 8-OH-DPAT. This may be relevant also to the diverse effects of 8-OH-DPAT on PPI in IR-sham and IR-depletion rats.

Finally, as IR downgraded the tissue concentration of 5HT in our data and others, IR rats, to a degree, appear equivalent to the 5-HT depleted SOC rats in terms of 5-HT-related performance, and thus the impact of 5,7-DHT in IR rats accordingly becomes less than that of SOC rats (Table 1). This hypothesis, together with an enhanced postsynaptic 5-HT1A activity of IR, may partially explain the observation that the effects of 8-OH-DAPT on PPI in SOC-depletion rats were similar to that of IR-sham IR rats.

In conclusion, the present study demonstrated a paradigm of using 5-HT depletion model in exploring the role of 5-HT1A receptor in the gating dysfunction induced by a lack of social connection since early life. The integrity of central 5-HT system is a key factor to modulate the sensorimotor gating only when the 5-HT1A is activated, in which the post-weaning social attachment is highly pertinent to the modulatory outcome.

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